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(54) Cyclic peptide antifungal agents and process for preparation thereof Zyklische Peptide mit antifungischer Wirkung und Verfahren zu ihrer Herstellung Dérivés cyclopeptidiques antifongiques et procédé de leur préparation

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Ba kground of the Invention

5 [0001] This invention relates to cyclic peptide antifungal agents. In particular, it relates to acyl derivatives of the echinocandin class of cyclic peptide antifungal agents; to methods for treating antifungal and parasitic infections, and to formulations useful in the methods.

[0002] The compounds provided by this invention are semi-synthetic antifungal agents in that they are derived from the cyclic peptide antifungals which are produced by culturing various microorganisms. A number of cyclic peptide antifungals are known. Among these are echinocandin B (A30912A), aculeacin, mulundocandin, sporiofungin, L-671,329, FR901379, and S31794/F1. All such antifungals are structurally characterized by a cyclic hexapeptide core, or nucleus, the amino group of one of the cyclic amino acids bearing a fatty acid acyl group forming a side chain off the core or nucleus. For example, echinocandin B has a linoleoyl side chain while aculeacin has a palmitoyl side chain. These fatty acid side chains of the cyclic hexa- peptides can be removed by enzymatic deacylation to provide the free nucleus. (Formula (1), hereinafter, wherein R₂ is hydrogen.) Reacylation of the amino group of the nucleus provides semisynthetic antifungal compounds. For example, the echinocandin B nucleus provides a number of antifungal agents when reacylated with certain unnatural side chain moieties (see Debono, U.S. Pat. No. 4,293,489). Among such antifungal compounds is cilofungin which is represented by the formula (1) wherein R is methyl, R₁ is hydrogen and R₂ is p-(n-octyloxy)benzoyl.

[0003] Enzymatic deacylation of the cyclic hexapeptides is carried out with deacylase produced by the organism Actinoplanes utahensis and related microorganisms as described by Abbott et al., U.S. Pat. No. 4,293,482.

[0004] The present invention provides acylated cyclic hexapeptides having unique side chain acyl groups which, inter alia impart enhanced antifungal and antiparasitic potency e.g. against pathogenic strains of Candida albicans. Also provided is a process for removing the aminal and benzylic hydroxy groups to result in a dideoxy compound of formula (1) (R = H).

Summary of the Invention

[0005] The compounds provided by this invention are represented by the following formula (1):

55 wherein

> R' is hydrogen, methyl or NH₂C(O)CH₂-; R" and R" are independently methyl or hydrogen;

R and Ry are independently hydroxy or hydrogen;

R₁ is hydroxy, hydrogen or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy; and

I) R₂ is a substituted benzoyl group represented by the formula

wherein

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R₃ is quinolyl; or

II) R₂ is an acyl group represented by the formula

wherein

Z is -C≡C-, -CH=CH-, or a carbon to carbon bond;

A) R_4 is C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, quinolyl or phenyl; or

B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkenyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, phenyl, substituted phenyl, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m and n are integers of from 2 to 4, and p is 0 or 1; or

- C) R₄ is phenyl substituted with C₁-C₆ alkoxy substituted by fluoro, bromo, chloro or iodo; or
- D) R₄ is a group represented by the formula

-Y-R₆

wherein

Y is -C≡C- or -C=C-; and

 R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} substituted alkyl; C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzothiazolyl, thienyl, phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(CH_2)_{p'}-W- R_5 wherein p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or piperazino, and R_5 is hydrogen, C_1 - C_{12} alkyl, C_3 - C_{12} cycloalkyl, benzyl or C_3 - C_{12} cycloalkylmethyl; or C_1 - C_6 alkoxy substituted by fluoro, bromo, iodo or chloro; or R_6 is a phenyl substituted by a polyoxa-alkyl group represented by the formula

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein m, n and p are as defined above; or R₂ is an acyl group represented by the formula

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Z is -C≡C- or -CH=CH-;

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A) R_4 is hydrogen, C_2 - C_{12} alkynyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkoxy; or B) R_4 is C_1 - C_{12} alkoxy substituted with C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_2 - C_{12} alkynyl, amino, C_1 - C_4 alkylamino, di- $(C_1$ - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino, phenyl substituted with a polyoxa-alkyl group represented by the formula

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$$\hbox{-O-(CH$_2)$_m$-[O-(CH$_2)$_n]$_p$-O-(C$_1$-C$_{12} alkyl)}$$

wherein m,n and p are as defined; or

C) R₄ is C₁-C₁₂ alkoxy substituted with a group of the formula

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wherein R_8 is C_1 - C_6 alkoxy optionally substituted with phenyl; or D) R_4 is a group represented by the formula

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wherein p', W and R_5 are as defined; or IV) R_2 is a group having the formula

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wherein Y and R₆ are as defined above; or V) R₂ is naphthoyl substituted with R₄

wherein

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A) R_4 is C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, quinolyl or phenyl; or

B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, phenyl, substituted phenyl, phenyl substituted with a polyoxa-alkyl group represented by the formula

wherein m, n and p are as defined; or

C) R₄ is phenyl substituted with C₁-C₆ alkoxy substituted by fluoro, bromo, chloro or iodo; or

D) R₄ is a group represented by the formula

30 wherein

Y has the same meanings as defined above; and

 R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} substituted alkyl; C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzothiazolyl, thienyl, phenyl substituted by amino, C_1 - C_{12} alkythio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(C_1), or C_1 - C_1 0 alkoxy substituted by fluoro, bromo, iodo or chloro; or

R₆ is a phenyl substituted by a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m, n and p are as defined above; and the pharmaceutically acceptable non-toxic salts thereof.

[0006] Also provided are formulations and methods for inhibiting parasitic and fungal activity which employ the compounds of the invention, and a process for preparing the dideoxy form of the compounds.

Detailed Description

[0007] The term: "C₁-C₁₂ alkyl" refers to the straight or branched chain alkyl hydrocarbon groups such as, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl, t-butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, undecyl and dodecyl groups; and the like.

[0008] The term "C₂-C₁₂ alkenyl" refers to groups such as vinyl, 1-propene-2-yl, 1-butene-4-yl, 1-pentene-5-yl, 1-butene-1-yl, and the like.

[0009] The term "C2-C12 alkynyl" refers to such groups as ethynyl, propynyl, pentynyl, butynyl and the like.

⁵⁵ [0010] The term "C₁-C₁₂ alkylthio" refers to such groups as methylthio, ethylthio, t-butylthio, and the like.

[0011] The term "C₁-C₁₂ alkoxy" refers to the straight or branched chain oxyalkyl groups such as, e.g. methoxy, ethoxy, propoxy, butoxy, heptoxy, octyloxy, dodecyloxy, and the like.

[0012] The term C₃-C₁₂ cycloalkoxy" refers to such groups as cyclopropoxy, cyclobutoxy and the like.

[0013] The term "C₃-C₁₂ cycloalkenyl" refers to such groups as cyclopropenyl, cyclobutenyl, cyclopentenyl, and the like.

[0014] The term " C_1 - C_{12} substituted alkyl," " C_2 - C_{12} substituted alkenyl", and " C_2 - C_{12} substituted alkynyl", denotes the above substituted one or two times with halogen, hydroxy, protected hydroxy, amino, protect d amino, C_1 - C_7 acyloxy, nitro, carboxy, protected carboxy, carbamoyl, carbamoyloxy, cyano, methylsulfonylamino, phenyl, substituted phenyl, or C_1 - C_{12} alkoxy.

[0015] The term "substituted phenyl" is represented by a phenyl group substituted with one, two, or three moieties chosen from halogen, hydroxy, protected hydroxy, cyano, nitro, C_1 - C_{12} alkyl, C_1 - C_{12} alkoxy, carboxy, protected carboxy, carboxymethyl, hydroxymethoyl, amino, aminomethyl trifluoromethyl or N-(methylsulfonylamino)

[0016] The term "C₃-C₁₂ cycloalkyl" refers to such groups as cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl and cycloheptyl.

[0017] The term "C₁-C₄ alkylamino" refers to such groups as methylamino, ethylamino, n-butylamino and the like.

[0018] The term "di-(C₁-C₄ alkyl)amino" refers to such groups as dimethylamino, diethylamino, di-n-propylamino, di-n-butylamino, methylethylamino, methyl-n-butylamino, and like tertiary amino groups.

[0019] The term "C₁-C₁₂ alkanoylamino" refers to such groups as acylamino groups derived from the C₁-C₁₂ carboxylic acids and are exemplified by formamido, acetylamino, propionylamino, butyrylamino, and the like,

[0020] The term " C_3 - C_{12} cycloalkylmethyl" refers to those C_3 - C_7 cycloalkyls described above further substituted by methyl.

[0021] The terms "C₇-C₁₀ bicycloalkyl" and "C₇-C₁₄ tricycloalkyl" refer to such groups as bicyclo[2.2.1.]hept-2-yl, bicyclo[2.2.1.]hep-4-en-2-yl, bicyclo[3.3.1.]nona-3-yl, bicyclo[3.3.1.]nona-2-yl, bicyclo[3.2.1.]oct-2-yl, bicyclo[2.2.2]oct-5-en-2-yl, adamantyl and the like.

[0022] The term "dideoxy" refers to compounds of the formula (1) wherein R=H.

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[0023] The term "inhibiting", such as used in relation to the methods for inhibiting parasitic and fungal activity, is defined to mean its normal definition, i.e., to stop, retard or prophylactically hinder or prevent.

[0024] The term "activity", as used in relation to parasitic and fungal activity, includes growth thereof and attending characteristics and results from the existence of the parasite or fungus.

[0025] The term "contacting", as used in relation to the methods for inhibiting parasitic and fungal activity by contacting a compound of the invention with a parasite or fungus, is defined to mean its normal definition. However, the term does not imply any further limitations to the process, such as by mechanism of inhibition, and the methods are defined to encompass the spirit of the invention, which is to inhibit parasitic and fungal activity by the action of the compounds and their inherent anti-parasitic and anti-fungal properties, or in other words, the compounds, used in the method are the causative agent for such inhibition.

[0026] Examples of acyl groups wherein R₂ is a group represented by the formula

are diphenyl acetylenes (Z=-C=C-), stilbenes (Z=-CH=CH-), and biphenyls (Z = a carbon to carbon bond). Among examples of such biphenyl groups, wherein Z is a carbon to carbon bond i.e. a phenyl to phenyl bond, are 4-{4-(butyloxy) phenyl]benzoyl, 4-{4-(cyclobutylmethoxy)-phenyl]benzoyl, 4-{4-cyclopentylmethoxy)phenyl]benzoyl, 4-{4-(cyclohexy-lethoxy)-phenyl]benzoyl, 4-phenylbenzoyl, 4-[4-(11-amino-undecyloxy)-phenyl]benzoyl, 4-{4-(11-formamidoundecyloxy)phenyl]benzoyl, 4-{4-(isopentyloxy)phenyl]benzoyl, and the like. Examples of diphenylacetylene and stilbene acyl groups, R₂, wherein Z is an acetylenic bond or an ethylene bond are 4-styrylbenzoyl, 4-(4-methoxystyryl)benzoyl, 4-(4-butyloxystyryl)benzoyl, 4-(phenylethynyl)benzoyl, 4-(4-ethoxyphenylethynyl) benzoyl, 4- (4-cyclohexyloxyphenylethynyl)benzoyl, and the like.

[0027] Examples of such acyl groups wherein R₄ is represented by the formula -Y-R₆ include 4-[4-(phenylethynyl) phenyl]benzoyl, 4-[4-(phenylethynyl)-phenoxy]benzoyl, 4-[4-(hexynyl)phenyl]benzoyl, 4-[4-(styryl)phenoxy]benzoyl, 4-[4-(4-benzylphenylethynyl)-phenyl]benzoyl, 4-[4-(4-methylpiperidino)ethoxy]phenylethynyl]phenyl]benzoyl and like acyl groups. Such acyl groups wherein R₄ is represented by the formula -O-(CH₂)_p-W-R₅ form salts of the basic amino groups of the piperidine and piperazine heterocyclic groups with both organic and inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid and phosphoric acid and with organic acids such as the sulfonic acids, benzenesulfonic acid, toluenesulfonic acid, methanesulfonic acid, acetic acid, chloroacetic acid, trifluoroacetic acid, benzoic acid, isophthalic acid, salicylic acid, citric acid, malic acid, succinic acid, malonic acid and like acids.

[0028] The following tables contain further examples of the cyclic peptides represented by the formula (1). Table 1 contains examples of cyclic peptides wherein the acyl group R₂ is of the formula

Table 1

<u>R2</u>

The following Table 2 illustrates the compound of the formula (1) wherein $\rm R_2$ is represented by the formula [0029]

Table 2

<u>R2</u>

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Table 2 continued R2

[0030] The acyl cyclohexapeptides represented by formula (1) exhibit antiparasitic activity, for example, they are especially active against the infectious fungi Candida albicans and Candida parapsilosis. They also exhibit significant activity against Aspergillus fumigatus. They are active both in vitro and in vivo and accordingly are useful in combating systemic fungal infections.

[0031] The compounds of the invention also inhibit the growth of certain organisms primarily responsible for opportunistic infections in immunosuppressed individuals. For example the compounds of the invention inhibit the growth of Pneumocystis carinii the causative organism of pneumocystis pneumonia in AIDS sufferers.

[0032] The antifungal activity of the compounds of the invention is determined in vitro in standard agar dilution tests and disc-diffusion tests wherein minimum inhibitory concentrations of the test compounds obtained. Standard in vivo tests in mice are used to determine the effective dose of the test compounds in controlling systemic fungal infections. [0033] Tables 4A-E below contain the minimum inhibitory concentrations (MIC) in micrograms per milliliter (mcg/ml) for compounds of the invention against Candida albicans and Candida parapsilosis, and for certain compounds, the effective dose, ED₅₀, in mice.

[0034] In Tables 4A-E, R'=CH₃, R"=CH₃, R"=CH₃, R"=OH, R₇=OH and R₁=H, In Tables 4A-D, R=OH, while in Table E, R=H.

[0035] In Table 4C, R2 is of the formula

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where Z is a carbon-carbon bond and R₄ is as indicated.

TABLE 4C

R ₄	MIC (mcg/ml)	ED ₅₀ (mg/ml)
	C.alb.	C.parap.	
-C≡C-C ₄ H ₉	0.039	2.5	1.20
-C≡C-C ₆ H ₅	0.039	0.625	0.60
-C ₆ H ₅	0.078	10	1.3

[0036] The non-dideoxy compounds of the invention (formula (1) are prepared with the amino nuclei of the cyclic hexapeptides which are represented by the formula when R_2 is hydrogen. These amino nuclei are obtained from the known natural products by the known enzymatic deacylation by which the fatty acid side chains of the natural compounds are removed. For example, echinocandin B which can be represented by the formula (1) wherein R'=R''=methyl, R is OH, Ry is hydroxy, R_1 is H, R_7 is OH, and R_2 is linoleoyl, is deacylated to provide the echinocandin B nucleus ($R_2=H$) with the deacylase produced by the organism Actinoplanes utahensis as described by U.S. Patent Nos. 4,293,482 and 4,304,716.

[0037] The known natural cyclic hexapeptides which are N-deacylated to provide the amino nuclei starting materials include echinocandin B (also known as A-30912A), aculeacin (palmitoyl side chain), tetrahydoechinocandin B (stearoyl side chain), mulundocandin (branched C₁₅ side chain), L-671,329 (C₁₆ branched side chain), S 31794/F1 (tetradecanoyl side chain), sporiofungin (C₁₅ branched side chain) and FR901379 (palmitoyl side chain). The amino nuclei obtained by the N-deacylation are then acylated by employing known amino acylation procedures to provide the N-

acyl cyclic hexapeptides represented by the formula (1) wherein R_2 represents the acyl groups defined hereinabove. The acylating moiety is preferably an active ester of the carboxylic acid RCOOH such as the 2,4,5-trichlorophenyl ester. The R_2 COOH precursor acids ar prepared by the hydrolysis of the nitrile R_2 CN or the ester R_2 COOC₁- R_2 COOC all. These nitrile and ester intermediates ar prepared by known methods.

- [0038] The alkoxy aromatic (ie. phenyl and biphenyl) compounds of Tables 9-10 are prepared by one of the two following procedures:
 - A. The hydroxyaromatic compound (1 equivalent) is dissolved in acetonitrile (200-300 ml) and a base, such as potassium t-butoxide or potassium carbonate,(1-equivalent), is added. An alkyl bromide, iodide, or p-toluenesulfonate (1 equivalent) is then added and the solution is refluxed for 6 hours. The solvent is evaporated in vacuo and the residue is dissolved in ether and 2N sodium hydroxide. The ether layer is dried over magnesium sulfate and evaporated to give the alkoxyaromatic product.
 - B. The hydroxyaromatic compound (1 equivalent), alkyl alcohol (1 equivalent), and triphenylphosphine (1 equivalent) are dissolved in tetrahydrofuran (200-300 ml) and diethylazodicarboxylate (1 equivalent) is added dropwise over 10 minutes at room temperature. After 17 hours the solvent is removed in vacuo and the residue is dissolved in ether. This organic layer is extracted with 2N sodium hydroxide solution, dried over magnesium sulfate, and evaporated to give a product which is crystallized from ether/pentane or, if the product contains a tertiary amine, the hydrochloride salt is formed and crystallized from methanol/ethyl acetate.

		Table 9		
			RO () ()	LOCH,
Alkylhalide or tosylate	Wt.	Method	& I	W.L.
I(CII ₂) ₂ CII ₃	2.6	A	-(CH ₂) ₂ CH ₃	4.4
H3C (SO3.(CH2)2O(CH2)3CH3	2.7	<	-(CH ₂) ₂ O(CH ₂) ₃ CH ₃	5.6
H3C (SO3.(CH2)20C(CH3)3	2.7	<	—(CH ₂) ₂ OC(CH ₃) ₃	2.6
		Table 10		
			Ro C	осн3
Alkylhalide or tosylate	Wt.	Method	₩	Wt
I(CI12)2CI13	3.8	A	-(CH ₂) ₂ CH ₃	1.4
H3C { _ } SO3.(СН2)2Q(СН2)3СН3	3.6	<	-(CH ₂) ₂ O(CH ₂) ₃ CH ₃	2.1
H3C \$ \$03.(CH2)20C(CH3)3	4.9	<	-(CH ₂) ₂ OC(CH ₃) ₃	5.2

[0039] The alkynyl and alkenyl aromatic compounds contained in Tables 11-14 are prepared by the following procedure:

[0040] An aromatic bromide, iodide, or trifluoromethane-sulfonate (1 equivalent) is dissolved in acetonitrile (600 ml/ 0.1 mole of aromatic reactant) under a nitrogen atmosphere. An alkyne or alkene (1 equivalent), triethylamine (2 equivalents), palladium dichloride (0.05 equivalents), triphenylphosphine (0.1 equivalents), and cuprous iodide (0.025 equivalents) are added and the solution is refluxed for 17 hours. The solvent is removed in vacuo and the residue is slurried in ether (300 ml). Solids are removed by filtration and the filtrate is washed with 1N hydrochloric acid solution. The organic layer is dried over magnesium sulfate and evaporated to yield the product.

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5			W S	26.2	28.1	1.9	11.2			₩ 80	2.6	5.1	23.3			a K	11.4
10		CH3		CH3 h (trans)	77		13)3	·	CCH ₃			-(CH ₂) ₃ CH ₃	-Si(CH ₃) ₃		CH3 CH3		(CH ₂) ₇ CH ₃
15		R D LOCHS	M	C== (CH2)2CH3	-C==-(CH ₂)/CH ₃	-C-	-Si(C)		Q	∞ 1)—————————————————————————————————————	-C==-(CH	-ESi(C		A COL		-C=-(CH
20	TABLE 11	·						TABLE 12	œ.	•	,			TABLE 13			
25		O LLOCH3	wt	28.8	28.8		11.5		- CH3	wt.	6.0	0.9	40.0		COCH,		2 - 3
30		Ç] ^{≱ -}	282	25	v .) }			4		Br		
35			W.L.	12.1	15.2	6.1	4.3			w k	æ. <u>–</u>	4.	10.9			W.L.	7.6
40			or olefin	(CH ₂) ₅ CH ₃	(CH ₂) ₂ CH ₃		-Si(CH ₃) ₃			ene		(CH ₂) ₃ CH ₃	(CH ₃) ₃			lene	-(CH ₂) ₇ CH ₃
45			Acetylene			111	N-E-Si			Acetylene) H	H==-(C	H==-SI(CH ₃),			Acetylene	(C)
50		1	·	1			1		1		i		1 1		1		•

5	W.E.	10.2	33
10	13	Choor, Coor,	2 — С — ОСН3
15	Product) C=C
20		40 P	
25	W.	34.4	1.2
30	-Pi	P P	
TABLE 14	Halide		
40	N N	10.5	1.2
45	ne	LOCH, LOCH,	ССН3
50	Acetylene		
55		± +	I

[0041] The aromatic boronic acids listed in Table 15 were prepared by the following procedure:

[0042] An aromatic halide (1 equivalent) is cooled to -78°C in tetrahydrofuran solvent. Butyl lithium (1.2 equivalents) is added. After 15 min triisopropyl borate (2 equivalents) is added and after 10 min of stirring the cooling bath is removed. When the reaction has warmed to room temperature water is added to quench the reaction followed by 1N HCI. The organic layer is removed under reduced pressure leaving a solid precipitate which is collected by filtration. This solid is washed with hexane leaving the pure boronic acid.

[0043] The terphenyl esters listed in Table 16 were made in the following manner:

[0044] An aromatic boronic acid (1 equivalent), methyl 4-iodobenzoate (1 equivalent), and potassium carbonate (1.5 equivalents) were mixed in a nitrogen-purged toluene solution. Alternatively, the trichloro phenyl ester of iodobenzoate my be used. Added tetrakis(triphenylphosphine)palladium (0.03 equivalents) and refluxed for 7 hrs. The solution was decanted to remove the potassium carbonate and reduced in vacuo. The residue was triturated with acetonitrile and the product solid was collected by filtration.

R=B(OII),	Wt. (g)	6.1	12.0	4.1	5.7	1.9		H,CO () () R	Wt. (g)	4.2	5.2	3.7	
TABLE 15 R=Br	Wt. (g)	_	31.0	10.9	13.6	5.0	TABLE 16] =	3.2	7. °C	3.6	•
		>O(CH ₂) ₃ CH ₃	O(CH ₂) ₄ CH ₃		O(CH ₂) ₂ O(CH ₂) ₃ CH ₃	O(CH ₂) ₂ OC(CH ₃) ₃		(HO) ₂ B-{	Wt. (g)	5.0	0.0 7.7	13 3.7	
		R-000	n \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	n () ()	RA SOLO	R 000	·	M		-0(CH ₂) ₃ CH ₃	-O(CH ₂),CH ₃	-0(CH ₂) ₂ O(CH ₂) ₃ CH ₃	ではいいのではいい。

[0045] The aromatic nitriles or carboxylate esters described in Tables 9-16 can be converted to carboxylic acids by

one of the two following hydrolysis procedures:

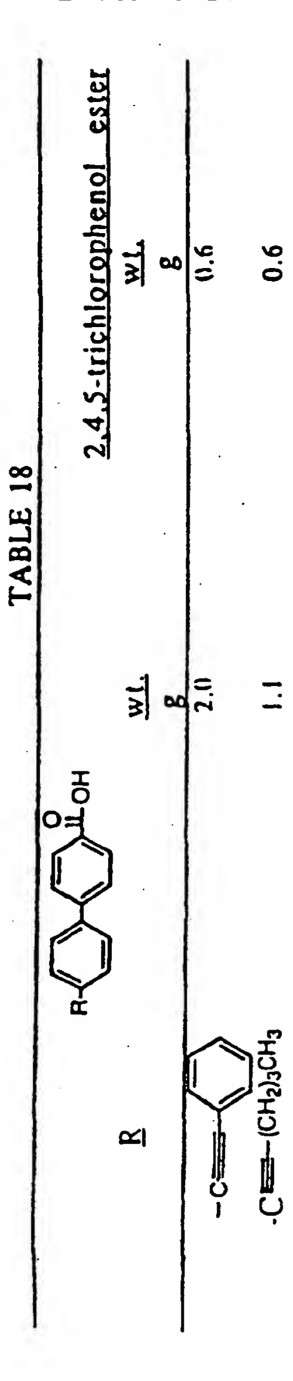
4.3

A. An aromatic nitrile is dissolved in ethanol and an excess of 50% sodium hydroxide solution and refluxed for 2 hours. Water is added until a solid precipitates. The precipitate is collected by filtration, added to dioxane and 6N hydrochloric acid solution and refluxed for 17 hours. Water is added and the carboxylic acid product crystallizes and is collected by filtration and dried under vacuum.

B. A carboxylate methyl ester is dissolved in methanol, excess 2N sodium hydroxide solution is added and the solution is refluxed for 5 hours. The solution is made acidic with excess hydrochloric acid and water is added until a precipitate forms. The carboxylic acid is collected by filtration and dried under vacuum.

[0046] The carboxylic acids are converted to 2,4,5-trichlorophenyl esters shown in Tables 18, 20 and 22-25 by the following general procedure:

[0047] The aromatic acid (1 equivalent), 2,4,5-trichlorophenol (1 equivalent), and N,N'-dicyclohexylcarbodiimide (1 equivalent) are dissolved in methylene chloride. The mixture is stirred for 17 hours after which it is filtered. The filtrate is evaporated to dryness and the residue is dissolved in ether, filtered, and pentane is added until crystallization begins. The crystalline product is collected by filtration and dried under vacuum.



5 10	E 20	.4.5-trichlo	1.5 E 22	2.4.5-trichlorophenol ester	1.2
25	TABLE 20	wt. 8	1.5 TABLE 22	WL.	0.8
30		п Дон		pia	· ·
40		≃1	<u></u>	Carboxylic ac	100 () () () () () () () () () (
<i>35 40</i>		R H		Carboxylic acid	

5	ester		ester		ester	
10	2,4,5-Trichlorophenol Wt. (g)	4.8 3.9 1.9	2,4,5-Trichlorophenol Wt. (g)	5.2 5.2 2.1	2,4,5-Trichlorophenol Wt. (g)	2.5 1.5 1.3
15	2,4,5-Tri		2,4,5-Tri		2,4,5-Tri	
20 FE 23	Œ		LE 24		SLE 25	
TAB 1		3.3 3.0 2.3 3.3 1.3	TABL	200	TAE	6.0
30	HO AL	E E Z E -	M M	6 4 4		2.2.2.2
35	I		, H		Ħ.	
40	씸	-0(CH ₂) ₃ CH ₃ -0(CH ₂) ₃ CH ₃ -0(CH ₂) ₅ CH ₃ -0(CH ₂) ₂ O(CH ₂) ₃ CH ₃	A	-0(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ O(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ OC(CH ₃) ₃	≃ i	-0(CH ₂) ₃ CH ₃ (CH ₂) ₂ O(CH ₂) ₃ CH ₃ O(CH ₂) ₂ OC(CH ₃) ₃
45		1 7 7 50		100-		100

[0048] The dideoxy compounds of formula (1) are prepared by removing the benzylic and aminal hydroxy groups. The process includes subjecting a non-dideoxy compound of formula (1) (wherein R₂ may be hydrogen or acyl) to a strong acid such as trichloroacetic acid, trifluoroacetic acid or borontrifluoride etherate with trifluoroacetic acid being preferred, and a reducing agent, such as sodium cyanoborohydride or triethylsilane, with triethylsilane being preferred. The reaction takes place at temperatures of between -5 and 70°C, and in a suitable solvent such as methylene chloride, chloroform or acetic acid, with dichloromethane being preferred. The acid should be present in an amount of 2 to 60 moles per mole of substrate, and the reducing agent should be present in an amount of 2 to 60 moles per mole of substrate. This process affords selective removal of the aminal and benzylic hydroxy groups.

[0049] The compounds represented by the formula (1) have improved properties over the previously known N-acyl

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hexapeptide antifungals. For example, in general the compounds exhibit oral bioavailability, a property which is important for any systemic antifungal agent. Also, numerous N-acyl compounds of the formula (1) have enhanced antifungal activity and enhanced water solubility.

[0050] Among the N-acyl hexapeptides represented by the formula (1) certain are preferred embodiments of the invention. The compounds wherein R_2 is a diphenyl acyl group

$$-C(0)$$

wherein Z is a carbon to carbon bond and R_4 is -Y- R_6 and R_6 is C_1 - C_{12} alkyl phenyl or substituted phenyl and Y is an acetylenic bond.

[0051] Examples of preferred compounds of the above mentioned group include compounds wherein R_4 is 4-[4-(phenylethynyl)-phenyl]benzoyl and 4-[4-(n-butylethynyl)phenyl]benzoyl.

[0052] Preferred cyclohexylpeptide compounds are represented by the formula 1 wherein R'=R''= methyl, R_1 is hydrogen and R_2 is a preferred acyl group as defined hereinabove.

[0053] Table 26 is a list of the most preferred R_2 substituents, wherein $R=R_7=R^y=OH$; $R'=R''=CH_3$; and $R_1=H$.

5			wt	80	26.2	9.0	28.1	1.9	11.2			W.	2.6	5.1	23.3			W a	4.11
10		CH2			CH	3 (trans)	Ç.		ele	•	o LOCH ₃			15 15	H ₃ l ₃		CCH,	•	JCH3
15		R D LOCHS	~		$C = -(CH_2)_5CH_3$	9	-CEEF-(CH2)7CH3)	- SI(CH			21	- C	-C==-(CH2)3CH3	-EE-SI(CH ₃) ₂		R KOLLO	M	-C==-(CH ₂),CH ₃
20	TABLE 11					-CH-	1			TABLE 12	H	•				TABLE 13			
25	Ţ	LOCH ₃			80	4	œ		5		- CH3	1 _1		a	0.		HOCH,	, 	
30		Y	W.	8	28.8	14.	28.8	e.	11.5				(1.0)	0.0	40.0		Br 40		=
<i>35</i>			WI.	80	12.1	6.1	15.2	1.9	4.3			w K	8. <u>-</u>	4.	6.01		•	* o	7.6
40			or olefin		(CH ₂) ₅ CH ₃	(CH ₂) ₅ CH ₃	(CH ₂) ₇ CH ₃	C	·Si(CH ₃) ₃			cue		-(CH ₂) ₃ CH ₃	-Si(CH ₃) ₃			cne	-(CH ₂) ₇ CH ₃
45			Acetylene		ı	Ì) - - -	Ī	HE			Acetylene	H	H-(C)	N-WH			Acetylene	H==-(Cl
				- 1	19				1 1				İ		1 1	1			1

5		t (g) FABMS	4 1142.4951**	0 1200.5336**	1194.5282*	9 1136.4832*	0 1194.5213*	4 1194.5247*	3 1126.5025*	1140.5103*	4 1154.5343*	5 1170.5234*	4 1170.5261*	2 1166.4758*	
15		Product	1.4	2.0	-	0.0	3.0	2.4	1.3	5.1	4.	6.5	1.4	0.2	
25		A30912	6.9	2.5	6.4	3.3	3.2	1.5	7.4	3.7	5.0	6.7	2.9	2.6	
30	TABLE 26	Ester	5.2	2.1	5.2	2.4	2.0	1.3	4.6	2.5	3.5	4.4	1.9	1.8	
35			3	9	9					બ	વ		100		
40 .		2													
45		R2	Н3С(СН3)26	CH ₂) ₂ O)(CH ₂) ₂ O	н ₃ С(СН ₃),0	(CH ₂) ₂ 0	CH ₂)20	H3C(CH2)3O	H3C(CH2),0	H ₃ C(CH ₂) ₅ O	%0 ² (2H2)0 ² (2H2)2 ² H	O(CH2)20		** m+ Li +
50			н3С(СН	(H ₃ C) ₃ CO(CH ₂) ₂ O	H3C(CH2)30(CH2)20	н3С(СН	H ₃ C(CH ₂) ₃ O(CH ₂) ₂ O	(H ₃ C) ₃ CO(CH ₂) ₂ O	H ₃ C(C	H ₃ C(H ₃ C((H³C(CH²)	(H3C)3CO(CH2)2		* m+1; **

⁵⁵ [0054] The N-acylhexapeptides provided by this invention are useful in the treatment of fungal infections both systemic infections and skin infections. Accordingly this invention also provides a method for treating fungal infections in man and animals which comprises administering to said host an antifungally effective non-toxic amount of an N-acyl-cyclohexapeptide represented by the formula 1. A preferred antifungal method comprises administering an N-acylhex-

apeptide compound where, in formula 1, R'=R''= methyl, R_1 is hydrogen and R_2 is a preferred acyl group as defined hereinabove.

[0055] The antifungal compound can be administered parenterally, e.g. i.m., i.p. or s.c., nasally, orally or can be applied topically for skin infections. The dose administered of course will vary depending on such factors as the nature and severity of the infection, the age and general health of the host and the tolerance of a particular host to the particular antifungal agent. The particular dose regimen likewise may vary according to such factors and may be given in a single daily dose or in multiple doses during the day. The regimen may last from about 2-3 days up to about 2-3 weeks or longer.

[0056] This invention also provides pharmaceutical formulations useful for administering the antifungal compounds of the invention. These formulations comprise an N-acylhexapeptide represented by the formula 1 or a pharmaceutically acceptable, non-toxic salt thereof and a pharmaceutically acceptable carrier.

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[0057] For parenteral administration the formulation comprises a compound of the formula 1 and a physiologically acceptable diluent such as deionized water, physiological saline, 5% dextrose and other commonly used diluents. The formulation may contain a solubilizing agent such as a polyethylene glycol or polypropylene glycol or other known solubilizing agent. Such formulations may be made up in sterile vials containing the antifungal and excipient in a dry powder or lyophilized powder form. Prior to use, the physiologically acceptable diluent is added and the solution withdrawn via syringe for administration to the patient. For oral administration, the antifungal compound is filled into gelatin capsules or formed into tablets. Such tablets also contain a binding agent, a dispersant or other suitable excipients suitable for preparing a proper size tablet for the dosage and particular antifungal compound of the formula 1. For pediatric or geriatric use the antifungal compound may be formulated into a flavored liquid suspension, solution or emulsion. A preferred oral carrier system is lineolic acid, cremophor RH-60 and water and preferably in the amount (by volume) of 8% lineolic acid, 5% cremophor RH-60, and 87% sterile water. The compound is added to the system in an amount of 2.5 to 40 mg/ml.

[0058] For topical use the antifungal compound can be formulated with a dry powder for application to the skin surface or it may be formulated in a liquid formulation comprising a solubilizing aqueous liquid or non-aqueous liquid, e.g., an alcohol or glycol. Such formulations are useful forms for use in the antifungal method provided herein.

[0059] The N-acylcyclohexapeptides provided herein may be formulated as described above in unit dosage formulations comprising for injection between about 50 mg and about 500 mg per vial. For oral use gelatin capsules or tablets comprising between about 100 mg and about 500 mg per capsule or tablet can be provided.

[0060] Preferred formulations of the invention comprises the active ingredient presented by the formula 1 wherein R'=R''= methyl, R_1 is hydrogen and R_2 is 4-[4-(phenylethynyl)-phenyl]benzoyl in gelatin capsules or as active ingredient the antifungal represented by the formula 1 wherein R'=R''= methyl, R_1 is hydrogen and R_2 is 4-[4-[2-(4-cyclohexyl-piperidino)ethoxy]phenyl]benzoyl or the hydrochloride salt form thereof in tablet or gelatin capsules. Further preferred formulations are those in which a preferred compound, as described above, is employed.

[0061] In yet a further aspect of the present invention there is provided a method for treating patients suffering from Pneumocystis pneumonia. The method can be used prophylactically to prevent the onset of the infection which is caused by the organism Pneumocystis carinii. The N-acylcyclicpeptide can be administered parenterally, e.g. via intramuscular (i.m), intravenous (iv.) or intraperitoneal (i.p.) injection, or orally or by inhalation directly into the airways of the lungs. Preferably the cyclic peptide is administered via inhalation of an aerosol spray formulation of the compound. [0062] An effective amount of a cyclic peptide will be between about 3 mg/kg of patient body weight to about 100 mg/kg. The amount administered may be in a single daily dose or multiple doses e.g. two, three or four times daily throughout the treatment regimen. The amount of the individual doses, the route of delivery, the frequency of dosing and the term of therapy will vary according to such factors as the intensity and extent of infection, the age and general health of the patient, the response of the patient to therapy and how well the patient tolerates the drug. It is known that PCP infections in AIDS patients are highly refractory owing to the nature of the infection. For example, in severe, advanced infections the lumenal surface of the air passages becomes clogged with infectious matter and extensive parasite development occurs in lung tissue. A patient with an advanced infection will accordingly require higher doses for longer periods of time. In contrast, immune deficient patients who are not severely infected and who are susceptible to PCP can be treated with lower and less frequent prophylactic doses.

[0063] The activity of the cyclicpeptide represented by the formula 1 is demonstrated in immunosuppressed rats. The tests were carried out in general as follows. One week after initiation of immunosuppression rats were inoculated intratracheally with parasites and maintained on immunosuppression for the remainder of the study. Prophylactic treatments began one day after parasite inoculation and therapeutic treatments began 3 or 4 weeks later after moderate PCP developed. Eight or ten animals were assigned to the following groups: those receiving test compound; non-treated Pneumocystis infected control animals; animals treated with trimethoprim-sulfamethoxazole (TMP-SMX); or non-treated, non-infected control animals. The efficacy of different treatments was evaluated by monitoring animal weights and survival during the studies and by determining the severity of PCP at necropsy. Stained impression smears of the lungs and stained lung homogenates were evaluated to determine the intensity of P. carinii infection.

[0064] The immune deficient rats employed in the tests were prepared as follows. Female Lewis rats weighing from

120-140 g each were immune suppressed with methyl prednisolone acetate at a dose of 4 mg/100 g for the first week, 3 mg/100 g for the second week and continuing weekly thereafter at 2 mg/100 g. All rats, except for the non-infected control rats, were inoculated intratracheally with 0.1 ml to 0.2 ml of Dulbecco's Modified Eagle Media containing between >10⁵ and 10⁶ P. carinii</sup> (trophozoites, precysts and cysts) harvested from the lungs of heavily infected donor animals (infection scores of 6) and maintained as cryopreserved (liquid nitrogen) inocula. Rats were maintained on immune suppression and PCP was allowed to develop for 3 or 4 weeks before initiation of therapy with test compounds. Body weights were recorded weekly and rats were allocated into treatment groups such that each group had a similar distribution of percent weight loss among animals. Rats were treated with test compounds for 2 or 3 weeks and then were necropsied. For prophylaxis studies, administration of test compound was initiated one day after intratracheal inoculation of parasites and was continued until the rats were necropsied.

[0065] Following the evaluation period for test compounds, the rats were necropsied and test results evaluated by Giemsa-stained, silver-methenamine stained impression smears and/or by silver-methenamine stained lung homogenate (see below). Necropsy was carried out as follows. The test rats were anesthetized with a mixture of ketamine hydrochloride and xylazine and then exsanguinated via the right atrium. Internal organs in the abdominal and thoracic cavities were examined for gross lesions.

[0066] A small portion of lung tissue from the left lobe of each rat was used to make the impression smears described below. Giemsa-stained impression smears were evaluated to determine the total number of parasites (trophozoites, precysts, and cysts). Impression smears from rats in groups whose treatments exhibited some anti-Pneumocystis activity (as judged by infection scores from Giemsa-stained slides) and from rats in the control groups were also stained with methamine silver, a stain specific for the cyst wall of the organism. Impression smears were randomized, numbered, and then evaluated. The infection scores used were as follows:

Score	Basis
0	No parasites found
1	1 to 5 parasites/10 oil fields
2	ca 1 parasite/field
3	2-10 parasites/field
4	>10 but <100 parasites/field
5	>100 but <1,000 parasites/field

A score of 6 was reserved for those infections with impression smears containing > 1,000 organisms/field (too numerous to count). Giemsa-stained slides were examined microscopically using a final magnification of 1008X. Methenamine silver-stained slides were examined with a final magnification of 400X.

[0067] Cysts in rat lung tissue were quantified as follows. A small portion of lung tissue from the left lobe of each rat was used to make impression smears as described above. The remainder of each lung was weighed, placed in a tube containing Hanks balanced salt solution (HBSS) (40X the lung weight) and homogenized using a Brinkman model tissue homogenizer. Two µl samples of the homogenized lung samples (1:4 dilution in HBSS) were placed in wells of teflon-coated, 12-well slides, stained with methenamine silver, and the number of cysts were scored as described above for the impression smears.

[0068] The activity and efficacy of a preferred N-acylcyclohexapeptide in the test animals is presented below. The compound of the formula 1 wherein R'=R"= methyl, R_1 is hydrogen and R_2 is 4[(4-phenylethynyl)phenyl]benzoyl when administered as an aerosol solution at a concentration of 5 mg/ml for one hour, twice weekly for 5 weeks resulted in 90% reduction in P carinii cysts in the lungs. When given orally at 10 mg/kg, bid for 3 weeks, the number of cysts in the lungs was reduced by >99% when compared with infected vehicle controls.

[0069] When the preferred N-acylcyclicpeptides were administered orally and by intraperitoneal injection the compound was effective in clearing P. carinii cysts from the lungs of heavily infected rats. For example, when the compound was administered at 10 or 40 mg/kg, bid for 4, 8 or 12 days, the number of identifiable cysts in the lungs of heavily infected rats was reduced by >99%. Similar efficacy was observed when the compound was administered i.p. at 1 mg/kg.

[0070] When tested orally for prophylactic activity, the preferred compound exhibited >99% cyst reduction in one of two studies when infected animals were dosed at 1 mg/kg and when given higher doses of 5 or 4 mg/kg.

[0071] The following examples of compounds of the invention and the manner of their preparation further describe the present invention.

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N-Acylation of Cyclohexp ptide Nuclei

[0072] The preparation of the d rivatives of the A30912A nucleus was accomplished by the following general procedure, with Table 27 listing these derivatives.

[0073] The A30912A nucleus and the 2,4,5-trichlorophenol ester are dissolved in dimethylformamide (25-50 ml) and stirred for 17-65 hours at room temperature. The solvent is removed *in vacuo* and the residue is slurried in ether and collected by filtration. The solid product is washed with methylene chloride and then dissolved in either methanol or acetonitrile/water (1:1 v/v). This solution is injected on a waters 600E semi-preparative chromatography system using a Rainin Dynamax-60A C₁₈ reverse-phase column. The column is eluted beginning with 20-40% aqueous acetonitrile and 0.5% monobasic ammonium phosphate (w/v) (monitored by UV at 230 nm and at a flow rate of 20 ml/min) until the unreacted A30912A nucleus is eluted and then deleting the buffer and eluting the product peak in aqueous acetonitrile. The fraction containing the product is evaporated in vacuo or lyophilized to provide the pure compound. The product may be analyzed by the same HPLC instrument using a Waters C₁₈ Micro Bondapak column and eluting with 40% aqueous acetonitrile containing 0.5% monobasic ammonium phosphate (w/v) at a 2 ml/min flow rate and monitoring the UV at 230 nm. The products may also be analyzed by fast atom bombardment mass spectrometry (FABMS). (In the compounds used, R'=R''=R'''=CH₃, R=OH, R₂=OH, R₁=H, R₇=OH, and R₂ is as defined).

5		HPLC Retention (min)	6.30	7.91	2.53
10		-	1078**	1058	1002 • •
15		Product (mg) FABMS	061	295	218
25	E 27 continued	A30912A Nucleus (g)	1.0	1.0	0.1
30	TABLE 27	Ester Reactant (mg)	968	571	108
35			여	역)
40		R2		C(CH ₂),	
45				ر (2	

5	HPLC Retention (min)	3.89
15	FABMS	1054**
20	Product (mg) FABMS	8
TABLE 27 continued	A30912A Nucleus (g)	1.0
TABLE	Ester Reactant (ing)	566 m+1Na1+
35		1
£0	R2	
15		TENIT TO

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[0074] Compounds such as those listed in Table 27 could be further modified at the phenolic hydroxy to provide R_7 = -OPO₃HNa as shown in Table 28. The procedure is as follows:

[0075] The lipopeptide (1 equivalent) and tetrabenzylpyrophosphate (2 equivalents) were dissolved in dimethylformamide which had been dried over 13X molecular sieves. Lithium hydroxide monohydrate (5 equivalents) was added and the stirred solution was monitored by HPLC. After 0.5 hr and 1 hr more lithium hydroxide (5 equivalents) was added. Between 1 and 2 hrs. the reaction was quenched with glacial acetic acid, the solvent removed under vacuum, and the residue purified over a semi-preparative C18 reverse-phase column using an aqueous acetonitrile eluent. The purified product was dissolved in (1/1) acetic acid/water with sodium acetate (1 equivalent) and 10% Pd/C catalyst.

The solution was placed under an atmosphere of hydrogen gas and stirred for 1 hr. After filtering to remove the catalyst, the solution was lyophilized to provide the pure final product. The purity was assessed by analytical HPLC and the product was analyzed by fast atom bombardment mass spectrometry (FABMS).

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10		FABMS	1228.4472*
15		Wt. (mg)	62
25	•	Prod.	-OPO ₃ HNa
30	TABLE 28	Wt. (mg)	300
<i>35</i> <i>40</i>		Start. Mat.	110-
45			000
50 55		R ₂	3C(CH ₂)3O

Preparation of dideoxy cyclohexapeptide

[0076] The preparation of the dideoxy compounds may be accomplished by the following proc dure.

[0077] To a suspension of a non-dideoxy cyclohexapeptide (formula (I) where R=OH and R₂ is hydrogen or acyl), in dichloromethane is added the reducing agent triethylsilane in dichloromethane. The solution is stirred and the volatile components are removed under reduced pressure and the residue triturated with diethyl ther. The compound is purified using HPLC, and the product lyophilized.

Example

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Dideoxycilofungin

[0078] To a suspension of cilofungin (10.00 g, 9.71 mmol) in dichloromethane (100 ml) was added a solution of triethylsilane (96 ml, 602 mmol) in dichloromethane (50 ml). Trifluoroacetic acid (46.4 ml, 602 mmol) was added as a solution in dichloromethane (50 ml) over 15 minutes. The solution was stirred at room temperature for two hours. The volatile reaction components were removed under reduced pressure and the residue triturated with diethyl ether. The compound was purified by reversed phase HPLC by means of a "Prep LC/System 500" unit (Waters Associates, Inc., Milford, Mass.) using a Prep Pak 500/C₁₈ Column (Waters Associates, Inc.) as the stationary phase. The column eluted with a gradient mobile phase using CH₃CN/H₂O (10:90 to 20:80 v/v) at 500 psi. The product containing fractions were pooled, evaporated under reduced pressure, and lyophilized from p-dioxane to yield dideoxycilofungin (6.66 g, 68.7%). FAB-MS: m/z calc. for C₄₉H₇₂N₇O₁₅, 998.5086; found, 998.512; UVλ(EtOH)nm(ε) 202.60(61012), 256.20(18569). [0079] A compound of the formula

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the preparation of which is discussed just prior to Table 27, can also be further modified at the phenolic hydroxy to provide R_7 =-OPO₃HNa, as indicated in the two paragraphs prior to Table 28. The compound produced is as follows:

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The product was analyzed by FABMS (using Lit) to give a peak at 1226.4853 (calculated for $C_{58}H_{74}N_7O_{20}$ PLi=1226.4886). Also, when analyzed by HPLC using a C18 reverse-phase column and eluting with 55% aqueous acetonitrile with 0.5% acetic acid at 2 ml/min and monitoring by UV at 280 nm, the compound had a retention time of 1.72 min.

Claims

1. A compound of the formula (1):

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E...

H. N. R. R. C. R. C. C. R. C. C. R. C. C. R.
wherein

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R' is hydrogen, methyl or NH₂C(O)CH₂-; R" and R" are independently methyl or hydrogen; R and R^y are independently hydroxy or hydrogen; R₁ is hydroxy, hydrogen or hydroxysulfonyloxy; R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy; and

I) R2 is a substituted benzoyl group represented by the formula

wherein R₃ is quinolyl; or (II) R₂ is an acyl group represented by the formula

- E- Tows - Tows.

wherein

Z is -C≡C-, -CH=CH-, or a carbon to carbon bond;

(A) R_4 is C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, quinolyl or phenyl; or

(B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkynyl, C_1 - C_1

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m and n are integers of from 2 to 4, and p is 0 or 1; or

C) R₄ is phenyl substituted with C₁-C₆ alkoxy substituted by fluoro, bromo, chloro or iodo; or

D) R₄ is a group represented by the formula

-Y-R₆

wherein

Y is -C≡C- or -C=C-; and

 R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} substituted alkyl; C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzothiazolyl, thienyl, phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(CH_2)_{p'}-W- R_5 wherein p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or piperazino, and R_5 is hydrogen, C_1 - C_{12} alkyl, C_3 - C_{12} cycloalkyl, benzyl or C_3 - C_{12} cycloalkylmethyl; or C_1 - C_6 alkoxy substituted by fluoro, bromo, iodo or chloro; or

R₆ is a phenyl substituted by a polyoxa-alkyl group represented by the formula

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein m, n and p are as defin d above; or R₂ is an acyl group represented by the formula

wherein

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Z is -C≡C- or -CH=CH-;

A) R_4 is hydrogen, C_2 - C_{12} alkynyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkoxy; or B) R_4 is C_1 - C_{12} alkoxy substituted with C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_2 - C_{12} alkynyl, amino, C_1 - C_4 alkylamino, di- $(C_1$ - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or C) R_4 is C_1 - C_{12} alkoxy substituted with a group of the formula

C || -NHCR.

wherein R_3 is C_1 - C_6 alkoxy optionally substituted with phenyl; or D) R_4 is a group represented by the formula

wherein p', W and R₅ are as defmed; or

IV) R₂ is a group having the formula

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wherein Y and R₆ are as defined above; or V) R₂ is naphthoyl substituted with R₄

wherein

A) R_4 is C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, quinolyl or phenyl; or

B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkenyl, C_2 - C_{12} substituted alkynyl, C_1 - C_1 -alkoxy, trifluoromethyl, phenyl, substituted phenyl, phenyl substituted with a polyoxa-alkyl group represented by the formula

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein m, n and p are as defined; or

C) R₄ is phenyl substituted with C₁-C₆ alkoxy substituted by fluoro, bromo, chloro or iodo; or

D) R₄ is a group represented by the formula

wherein Y has the same meanings as defined above; and

 R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} substituted alkyl; C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzothiazolyl, thienyl, phenyl substituted by amino, C_1 - C_{12} alkythio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(C_1 - C_2), or C_1 - C_2 alkoxy substituted by fluoro, bromo, iodo or chloro; or

R₆ is a phenyl substituted by a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m, n and p are as defined above; and the pharmaceutically acceptable non-toxic salts thereof.

- 2. A compound according to claim 1 wherein R₁ is hydroxy or hydrogen and R₇ is hydroxy or hydrogen.
- 3. A compound according to claim 1 or claim 2 wherein R', R" and R" are methyl, R₁ is hydrogen, and R₇ and R⁹ are OH.
 - 4. A compound according to any one of claims 1 to 3 wherein R2 is of the formula

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Z is a carbon to carbon bond; and

R₄ is C₃-C₇ cycloalkoxy; or

R₄ is phenyl substituted by C₁-C₁₂ alkoxy or phenyl substituted with a polyoxa-alkyl group of the formula

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wherein

 R_4 is a group of the formula -Y- R_6 , wherein Y is -C=C- or -C=C- and R_6 is C_1 - C_6 alkyl, phenyl, or phenyl substituted with a polyoxa-alkyl group of the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl}).$$

5. A compound according to any one of claims 1 to 3 wherein R₂ is of the formula

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Z is -C≡C-; and

R₄ is phenyl substituted by C₁-C₁₂ alkoxy or phenyl substituted with a polyoxa-alkyl group of the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

or

R₄ is a group of the formula

$$\text{-O-(CH}_2)_{p'}\text{-W-R}_5$$

wherein W is a piperidine group.

- 6. A compound according to any one of claims 1 to 3 wherein R is hydrogen.
- 7. A compound according to any one of claims 1 to 3 wherein R₂ is 4-[4-(phenylethynyl)phenyl]benzoyl or 4-[4-(n-butylethynyl)phenyl]benzoyl.
- 8. A compound of the formula (1):

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wherein

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R' is hydrogen, methyl or $NH_2C(O)CH_2$ -;

R" and R" are independently methyl or hydrogen;

R and Ry are independently hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy; and

I) R₂ is a group of the formula

and pharmaceutically acceptable salts thereof.

- 9. A compound according to claim 8 wherein R', R" and R" are methyl, R₁ is hydrogen and R₇ and R^y are hydroxy.
- 10. A compound of the formula (1):

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wherein

R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" is methyl or hydrogen;

R is hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy;

R₂ is an acyl group represented by the formula

wherein Z is -C≡C-, -CH=CH-, or a carbon to carbon bond;

 R_4 is C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, phenyl substituted by amino, C_1 - C_{12} alkythio, halogen, C_1 - C_{12} alkyl, C_1 - C_{12} alkoxy, trifluoromethyl, phenyl, or C_1 - C_6 alkoxy substituted by fluoro, bromo, chloro or iodo;

or R₄ is C₃-C₁₂ cycloalkoxy;

or R_4 is a group represented by the formula -Y-R₆ wherein Y is -C=C- or -CH=CH- and R₆ is C₁-C₁₂ alkyl, C₁-C₁₂ alkyl substituted by phenyl; C₃-C₁₂ cycloalkyl, phenyl, C₃-C₁₂ cycloalkenyl, naphthyl, benzthiazol-2-yl, or phenyl substituted by amino, C₁-C₁₂ alkythio, halogen, C₁-C₁₂ alkyl, C₁-C₁₂ alkenyl. C₁-C₁₂ alkoxy, trifluoromethyl, -O-(CH₂)_{p'}-W-R₅ wherein p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or piperazino, and R₅ is hydrogen, C₁-C₁₂ alkyl, C₃-C₁₂ cycloalkyl, benzyl or C₃-C₁₂ cycloalkylmethyl; or C₁-C₆ alkoxy substituted by fluoro, bromo, iodo or chloro;

or R₂ is an acyl group represented by the formula

wherein

Z is -C≡C- or -CH=CH-;

R₄ is hydrogen:

 R_4 is C_1 - C_{12} alkoxy, C_1 - C_{12} alkoxy substituted by C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, amino, C_1 - C_4 alkylamino, di- $(C_1$ - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino or a group of the formula

O || -NHCR:

wherein R₈ is C₁-C₆ alkoxy optionally substituted with phenyl; or R₄ is a group represented by the formula

25 or

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R₂ is a group selected from

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wherein

Y and R₆ are as defined above; or

R₂ is naphthoyl substituted with R₄ wherein

 R_4 is C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, phenyl substituted by amino, C_1 - C_{12} alkythio, halogen, C_1 - C_{12} alkyl, C_1 - C_{12} alkoxy, trifluoromethyl, phenyl, or C_1 - C_6 alkoxy substituted by fluoro, bromo, chloro or iodo;

or R₄ is C₃-C₁₂ cycloalkoxy;

or R_4 is a group represented by the formula -Y- R_6 wherein Y has the same meanings as defined above and R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} alkyl substituted by phenyl; C_3 - C_{12} cycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzthiazol-2-yl, or phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_1 - C_{12} alkenyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(C_1), or C_1 - C_2 alkoxy substituted by fluoro, bromo, iodo or chloro; and the pharmaceutically acceptable non-toxic salts thereof.

- 11. A compound according to claim 10 wherein R₁ is not hydroxysulfonyloxy and R₇ is not hydroxysulfonyloxy or phosphonooxy.
- 12. A compound of the formula

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13. A compound of the formula

14. A compound of the formula

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 $\text{wherein R is -O(CH}_2)_3\text{CH}_3, \ -\text{O(CH}_2)_4\text{CH}_3, \ -\text{O(CH}_2)_5\text{CH}_3, \ -\text{O(CH}_2)_2\text{O(CH}_2)_3\text{CH}_3 \ \text{or -O(CH}_2)_2\text{OC(CH}_3)_3. \\$

- 15. A compound according to claim 14 wherein R is -O(CH₂)₄CH₃.
- 16. A compound according to any of claims 1-15 for use in inhibiting parasitic activity.
 - 17. A compound according to any one of claims 1-15 for use in inhibiting fungal activity.
- 18. A compound according to any of claims 1-15 for use in inhibiting the growth of organisms responsible for opportunistic infections in immunosuppressed individuals.
 - 19. A compound according to any of claims 1-15 for use in inhibiting the growth of Pneumocystis carinii.

- 20. A pharmaceutical formulation comprising a compound according to any of claims 1-15 and a suitable pharmaceutical carrier.
- 21. A process for the preparation of a compound of the formula (1):

wherein

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R' is hydrogen, methyl or $NH_2C(O)CH_2$ -;

R" and R" is methyl or hydrogen;

R is hydrogen;

Ry is hydroxy or hydrogen;

R₁ is hydroxy, or hydrogen;

R₇ is hydroxy, or hydrogen; and

R₂ is hydrogen or acyl;

comprising the step of subjecting a compound of formula (1) wherein R=OH, to a strong acid in the presence of a reducing agent, in a suitable solvent.

22. A process for producing an N-acyl cyclic hexapeptide which process comprises acylating an amino nucleus of Echinocandin B with an active ester of a carboxylic acid represented by the formula

wherein R is $-O(CH_2)_3CH_3$, $-O(CH_2)_4CH_3$, $-O(CH_2)_5CH_3$, $-O(CH_2)_2O(CH_2)_3CH_3$ or $-O(CH_2)_2OC(CH_3)_3$.

23. A process for preparing a compound of the formula

wherein R is $-O(CH_2)_3CH_3$, $-O(CH_2)_4CH_3$, $-O(CH_2)_5CH_3$, $-O(CH_2)_2O(CH_2)_3CH_3$ or $-O(CH_2)_2OC(CH_3)_3$ which process comprises acylating an amino nucleus of Echinocandin B with an active ester of a carboxylic acid represented by the formula

24. A process according to claim 22 or claim 23 wherein R is

- 25. A process according to any one of claims 22-24 wherein the active ester is a 2,4,5-trichlorophenyl ester.
- 26. A process according to any one of claims 22-25 wherein the amine nucleus of Echinocandin B is obtained by N-deacylation of a naturally occurring cyclic hexapeptide.
 - 27. A process according to claims 26 wherein the naturally occurring cyclic hexapeptide is echinocandin B, tetrahydroechinocandin B, mulundocandin, L-671 329, S 31794/FI, sporiofungin or FR 901379.

Patentansprüche

1. Verbindungen der Formel (1):

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worin

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R' Wasserstoff, Methyl oder NH₂C(O)CH₂- ist;

R" und R" unabhängig voneinander Methyl oder Wasserstoff sind;

R und Ry unabhängig voneinander Hydroxy oder Wasserstoff sind;

R₁ Hydroxy, Wasserstoff oder Hydroxysulfonyloxy ist;

R7 Hydroxy, Wasserstoff, Hydroxysulfonyloxy oder Phosphonooxy ist; und

I) R₂ eine substituierte Benzoylgruppe der Formel

ist, worin R₃ Chinolyl ist; oder

II) R₂ eine Acylgruppe der Formel

ist, worin

Z -C=C-; -CH=CH-, oder eine Kohlenstoff-zu-Kohlenstoff-Bindung ist;

- (A) R_4 C_3 - C_{12} -Cycloalkyl, C_7 - C_{10} -Bicycloalkyl, C_7 - C_{14} Tricycloalkyl, C_3 - C_{12} -Cycloalkoxy, Naphthyl, Pyridyl, Thienyl, Benzothienyl, Chinolyl oder Phenyl ist; oder
- (B) R_4 Phenyl, substituiert durch Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_2 - C_{12} -Alkenyl, C_2 - C_{12} -Alkynyl, substituiertes C_1 - C_{12} -Alkyl, substituiertes C_2 - C_{12} -Alkenyl, substituiertes C_2 - C_{12} -Alkynyl, C_1 - C_1 -Alkoxy, Trifluormethyl, Phenyl, substituiertes Phenyl, Phenyl, substituiert mit einer Polyoxa-Alkylgruppe

der Formel

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worin m und n ganze Zahlen von 2 bis 4 sind und p 0 oder 1 ist; oder

(C) R₄ Phenyl, substituiert mit C₁-C₆-Alkoxy, substituiert durch Fluor, Brom, Chlor oder lod, ist; oder

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(D) R₄ eine Gruppe der Formel

-Y-R_e

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ist, worin

Y -C≡C- oder -C=C-, ist; und

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 R_6C_1 - C_{12} -Alkyl, substituiertes C_1 - C_{12} -Alkyl; C_3 - C_{12} -Cycloalkyl, C_7 - C_{10} -Bicycloalkyl, C_7 - C_{14} -Tricycloalkyl, Phenyl, C_3 - C_{12} -Cycloalkenyl, Naphthyl, Benzothiazolyl, Thienyl, Phenyl, substituiert durch Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_2 - C_{12} -Alkenyl, C_2 - C_{12} -Alkynyl, C_1 - C_{12} -Alkoxy, Trifluormethyl, -O-(CH_2) $_p$ -W- R_5 , worin p eine ganze Zahl von 2 bis 4 ist, W Pyrrolidino, Piperidino oder Piperazino ist, ist, und R_5 Wasserstoff, C_1 - C_{12} -Alkyl, C_3 - C_{12} -Cycloalkyl, Benzyl oder C_3 - C_{12} -Cycloalkylmethyl ist; oder C_1 - C_6 -Alkoxy, substituiert durch Fluor, Brom, lod oder Chlor, ist; oder

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R₆ ein Phenyl, substituiert durch eine Polyoxa-Alkylgruppe der Formel

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12}-A!kyl)$$

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ist, worin m, n und p wie oben definiert sind, oder

R₂ eine Acylgruppe der Formel

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ist, worin

Z -C=C- oder -CH=CH- ist;

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A) R_4 Wasserstoff, C_2 - C_{12} -Alkynyl, substituiertes C_2 - C_{12} -Alkynyl, C_1 - C_{12} -Alkoxy ist; oder

B) R₄ C₁-C₁₂-Alkoxy, substituiert durch C₃-C₁₂-Cycloalkyl, C₇-C₁₀-Bicycloalkyl, C₇-C₁₄-Tricycloalkyl, C₂-C₁₂-Alkynyl, Amino, C₁-C₄-Alkylamino, Di-(C₁-C₄alkyl)amino, C₁-C₁₂-Alkanoylamino, Phenyl, substituiert mit einer Polyoxa-Alkylgruppe der Formel

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worin m, n und p wie definiert sind, ist; oder

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C) R₄ C₁-C₁₂-Alkoxy, substituiert mit einer Gruppe der Formel

O || -NHCR₈

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worin R₈ C₁-C₆-Alkoxy, gegebenenfalls substituiert mit Phenyl ist, ist; oder

D) R₄ eine Gruppe der Formel

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$$-O-(CH_2)_{p'}-W-R_5$$

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ist, worin p', W und R_5 wie definiert sind; oder

IV) R₂ eine Gruppe der Formel

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- S - Y - E

ist, worin Y und R₆ wie oben definiert sind; oder

V) R₂ Naphthoyl ist, substituiert mit R₄, worin

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(A) R₄ C₃-C₁₂-Cycloalkyl, C₇-C₁₀-Bicycloalkyl, C₇-C₁₄ Tricycloalkyl, C₃-C₁₂-Cycloalkoxy, Naphthyl, Pyridyl, Thienyl, Benzothienyl, Chinolyl oder Phenyl ist; oder

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(B) R_4 Phenyl, substituiert durch Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_2 - C_{12} -Alkynyl, substituiertes C_1 - C_{12} -Alkyl, substituiertes C_2 - C_{12} -Alkenyl, substituiertes C_2 - C_{12} -Alkynyl, C_1 - C_{12} -Alkoxy, Trifluormethyl, Phenyl, substituiertes Phenyl, Phenyl, substituiert mit einer Polyoxa-Alkylgruppe der Formel

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12}-Alkyl),$$

worin m, n und p wie definiert sind, ist; oder

(C) R₄ Phenyl, substituiert mit C₁-C₆-Alkoxy, substituiert durch Fluor, Brom, Chlor oder lod, ist; oder

(D) R₄ eine Gruppe der Formel

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-Y-R₆

ist, worin

Y die gleichen Bedeutungen wie oben definiert besitzt; und

 R_6C_1 - C_{12} -Alkyl, substituiertes C_1 - C_{12} -Alkyl; C_3 - C_{12} -Cycloalkyl, C_7 - C_{10} -Bicycloalkyl, C_7 - C_{14} -Tricycloalkyl, Phenyl, C_3 - C_{12} -Cycloalkenyl, Naphthyl, Benzothiazolyl, Thienyl, Phenyl, substituiert durch Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_2 - C_{12} -Alkenyl, C_2 - C_{12} -Alkynyl, C_1 - C_1 -Alkoxy, Trifluormethyl, -O-(CH_2)_{p'}-W- R_5 , oder C_1 - C_6 -Alkoxy, substituiert durch Fluor, Brom, lod oder Chlor, ist; oder

Re ein Phenyl, substituiert durch eine Polyoxa-Alkylgruppe der Formel

-O-(CH₂)_m-[O-(CH₂)_n]_p-O-(C₁-C₁₂-Alkyl)

ist, worin m, n und p wie oben definiert sind; und pharmazeutisch zulässige nichttoxische Salze davon.

- 2. Verbindung nach Anspruch 1, worin R₁ Hydroxy oder Wasserstoff ist und R₇ Hydroxy oder Wasserstoff ist.
- 3. Verbindung gemäß Anspruch 1 oder Anspruch 2, worin R', R" und R" Methyl sind, R₁ Wasserstoff ist und R₇ und R^y OH sind.
- 4. Verbindung gemäß mindestens einem der Ansprüche 1 bis 3, worin R₂ von folgender Formel ist

-c(c)

worin

Z eine Kohlenstoff-zu-Kohlenstoff-Bindung ist; und

R₄ C₃-C₇-Cycloalkoxy ist; oder

R₄ Phenyl, substituiert durch C₁-C₁₂-Alkoxy, oder Phenyl, substituiert mit einer Polyoxa-Alkylgruppe der Formel

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12}-Aikyl),$$

ist; oder

 R_4 eine Gruppe der Formel -Y- R_6 ist, worin Y -C=C- oder -C=C- ist, und R_6 C_1 - C_6 -Alkyl, Phenyl oder Phenyl, substituiert mit einer Polyoxa-Alkylgruppe der Formel

-O-(CH₂)_m-[O-(CH₂)_n]_p-O-(C₁-C₁₂-Alkyl),

ist.

5. Verbindung gemäß mindestens einem der Ansprüche 1 bis 3, worin R2 die Formel

aufweist, worin

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Z -C≡C- ist; und

R₄ Phenyl, substituiert durch C₁-C₁₂-Alkoxy- oder Phenyl, substituiert mit einer Polyoxa-Alkylgruppe der Formel

ist, oder R₄ eine Gruppe der Formel

-O-(CH₂)_{p'}-W-R₅

ist, worin W eine Piperidingruppe ist.

- 6. Verbindung gemäß mindestens einem der Ansprüche 1 bis 3, worin R Wasserstoff ist.
- 7. Verbindung gemäß mindestens einem der Ansprüche 1 bis 3, worin R₂ 4-[4-(Phenylethynyl)phenyl]benzoyl oder
 4-[4-(n-Butylethynyl)phenyl]benzoyl ist.
 - 8. Verbindung der Formel (1):

worin

R' Wasserstoff, Methyl oder NH₂C(O)CH₂- ist;

R" und R" unabhängig voneinander Methyl oder Wasserstoff sind; R und R^y unabhängig voneinander Hydroxy oder Wasserstoff sind; R₁ Hydroxy, Wasserstoff oder Hydroxysulfonyloxy ist; R₇ Hydroxy, Wasserstoff, Hydroxysulfonyloxy oder Phosphonooxy ist; und

I) R₂ eine Gruppe der Formel

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H₂C(CH₂)₂O

H₃C(CH₂)₂O

H₃C(CH₂)₃O(CH₂)₂O

H₃C(CH₂)₃O(CH₂)₂O

H₃C(CH₂)₃O(CH₂)₂O

C

H₃C(CH₂)₃O(CH₂)₂O

C

H₃C(CH₂)₃O(CH₂)₂O

C

H₃C(CH₂)₃O(CH₂)₂O

(H₃C)₃C(CH₂)₃O(CH₂)₂O

ist, und pharmazeutisch annehmbare Salze davon.

- 9. Verbindung gemäß Anspruch 8, worin R', R" und R" Methyl sind, R₁ Wasserstoff ist und R₇ und R^y Hydroxy sind.
- 10. Verbindung der Formel (1):

worin

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25 R' Wasserstoff, Methyl oder NH₂C(O)CH₂- ist;

R" Methyl oder Wasserstoff ist;

R Hydroxy oder Wasserstoff ist;

R₁ Hydroxy, Wasserstoff oder Hydroxysulfonyloxy ist;

R7 Hydroxy, Wasserstoff, Hydroxysulfonyloxy oder Phosphonooxy ist;

R₂ eine Acylgruppe der Formel

ist, worin Z -C≡C-, -CH=CH- oder eine Kohlenstoff-zu-Kohlenstoff-Bindung ist;

R₄ C₃-C₁₂-Cycloalkyl, C₇-C₁₀-Bicycloalkyl, C₇-C₁₄-Tricycloalkyl, Phenyl, Phenyl, substituiert durch Amino, C₁-C₁₂-Alkylthio, Halogen, C₁-C₁₂-Alkyl, C₁-C₁₂-Alkoxy, Trifluormethyl, Phenyl oder C₁-C₆-Alkoxy, substituiert durch Fluor, Brom, Chlor oder lod, ist;

oder R₄ C₃-C₁₂-Cycloalkoxy ist;

oder R_4 eine Gruppe der Formel -Y- R_6 ist, worin Y -C \equiv C- oder -CH \equiv CH- ist, und R_6 C_1 - C_{12} -Alkyl, C_1 - C_{12} -Alkyl, substituiert durch Phenyl; C_3 - C_{12} -Cycloalkyl, Phenyl, C_3 - C_{12} -Cycloalkenyl, Naphthyl, Benzthiazol-2-yl oder Phenyl, substituiert durch Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_1 - C_{12} -Alkenyl, C_1 - C_{12} -Alkenyl, C_1 - C_1 -Alkyl, C_1 - C_1 -Alkyl, C_1 - C_1 -Alkyl, C_1 -Cycloalkyl, Pyrrolidino, Piperidino oder Piperazino ist, ist, und C_1 - C_1 -Alkyl, C_1 - C_1 -Alkyl, C_2 - C_1 -Cycloalkyl, Benzyl oder C_3 - C_1 -Cycloalkylmethyl ist; oder C_1 - C_6 -Alkoxy, substituiert durch Fluor, Brom, lod oder Chlor, ist;

oder R₂ eine Acylgruppe der Formel

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ist, worin Z -C≡C- oder -CH=CH- ist;

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R₄ Wasserstoff ist;

 $R_4\ C_1-C_{12}-Alkoxy,\ C_1-C_{12}-Alkoxy,\ substituiert\ durch\ C_3-C_{12}-Cycloalkyl,\ C_7-C_{10}-Bicycloalkyl,\ C_7-C_{14}-Tricycloalkyl,\ Amino,\ C_1-C_4-Alkylamino,\ Di-(C_1-C_4-alkyl)amino,\ C_1-C_{12}-Alkanoylamino\ oder\ eine\ Gruppe\ der\ Formel$

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worin R₈ C₁-C₆-Alkoxy, optional substituiert mit Phenyl, ist; oder R₄ eine Gruppe der Formel

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ist, oder

R₂ eine Gruppe ist, die aus

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gewählt ist, worin Y und R₆ wie oben definiert sind; oder

R₂ Naphthyl, substituiert mit R₄ ist, worin

 R_4 C_3 - C_{12} -Cycloalkyl, C_7 - C_{10} -Bicycloalkyl, C_7 - C_{14} -Tricycloalkyl, Phenyl, Phenyl, substituiert mit Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_1 - C_{12} -Alkoxy, Trifluormethyl, Phenyl oder C_1 - C_6 -Alkoxy, substituiert durch Fluor, Brom, Chlor oder lod, ist;

oder R₄ C₃-C₁₂-Cycloalkoxy ist;

oder R₄ eine Gruppe der Formel -Y-R₆ ist, worin Y die gleiche Bedeutung wie obenstehend besitzt, und R₆

 C_1 - C_{12} -Alkyl, C_1 - C_{12} -Alkyl, substituiert durch Phenyl; C_3 - C_{12} -Cycloalkyl, Phenyl, C_3 - C_{12} -Cycloalkenyl, Naphthyl, Benzthiazol-2-yl oder Phenyl, substituiert durch Amino, C_1 - C_{12} -Alkylthio, Halogen, C_1 - C_{12} -Alkyl, C_1 - C_{12} -Alkoxy, Trifluormethyl, -O-(CH_2) $_p$ -W-R $_5$ oder C_1 - C_6 -Alkoxy, substituiert durch Fluor, Brom, lod oder Chlor, ist; und die pharmazeutisch annehmbaren nicht-toxischen Salze davon.

- 11. Verbindung gemäß Anspruch 10, worin R₁ nicht Hydroxysulfonyloxy ist und R₇ nicht Hydroxysulfonyloxy oder Phosphonooxy ist.
- 12. Verbindung der Formel

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13. Verbindung der Formel

14. Verbindung der Formel

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25 НÔ OH H₃C *30* 0 HN HO CH₃ H₃C OH HK 35 HO OH 40

worin R -O(CH₂)₃CH₃, -O(CH₂)₄CH₃, -O(CH₂)₅CH₃, -O(CH₂)₂O(CH₂)₃CH₃ oder -O(CH₂)₂OC(CH₃)₃ ist.

- 15. Verbindung gemäß Anspruch 14, worin R -O(CH₂)₄CH₃ ist.
- 16. Verbindung gemäß mindestens einem der Ansprüche 1-15 zur Verwendung bei der Inhibierung parasitischer Aktivität.
 - 17. Verbindung gemäß mindestens einem der Ansprüche 1-15 zur Verwendung bei der Inhibierung fungaler Aktivität.
 - 18. Verbindung gemäß mindestens einem der Ansprüche 1-15 zur Verwendung bei der Inhibierung des Wachstums von Organismen, die für opportunistische Infektionen bei Individuen mit Immunosuppression verantwortlich sind.
 - 19. Verbindung gemäß mindestens einem der Ansprüche 1-15 zur Verwendung bei der Inhibierung des Wachstums von *Pneumocystis carinii*.

- 20. Pharmazeutische Formulierung, umfassend eine Verbindung gemäß mindestens einem der Ansprüche 1-15 und einen geeigneten pharmazeutischen Träger.
- 21. Verfahren zur Herstellung einer Verbindung der Formel (1):

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R' Wasserstoff, Methyl oder NH₂C(O)CH₂- ist;

R" und R" Methyl oder Wasserstoff ist;

R Wasserstoff ist;

Ry Hydroxy oder Wasserstoff ist;

R₁ Hydroxy oder Wasserstoff ist;

R7 Hydroxy oder Wasserstoff ist; und

R₂ Wasserstoff oder Acyl ist;

umfassend die Schritte des Unterziehens einer Verbindung der Formel (1), worin R=OH ist, einer starken Säure in Gegenwart eines Reduktionsmittels in einem geeigneten Lösungsmittel.

22. Verfahren zur Herstellung von cyclischem N-Acyl-Hexapeptid, wobei das Verfahren das Acylieren eines Aminokerns von Echinocandin B mit einem aktiven Ester einer Carbonsäure der Formel

umfasst, worin R -O(CH₂)₃CH₃, -O(CH₂)₄CH₃, -O(CH₂)₅CH₃, -O(CH₂)₂O(CH₂)₃CH₃ oder -O(CH₂)₂OC(CH₃)₃ ist.

23. Verfahren zur Herstellung einer Verbindung der Formel

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worin R -O(CH₂)₃CH₃, -O(CH₂)₄CH₃, -O(CH₂)₅CH₃, -O(CH₂)₂O(CH₂)₃CH₃ oder -O(CH₂)₂OC(CH₃)₃ ist, wobei das Verfahren die Acylierung eines Aminokerns von Echinocandin B mit einem aktiven Ester einer Carbonsäure der Formel

umfasst.

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- 24. Verfahren gemäß Anspruch 22 oder Anspruch 23, worin R -O(CH₂)₄CH₃ ist.
- 25. Verfahren gemäß mindestens einem der Ansprüche 22-24, bei dem der aktive Ester ein 2,4,5-Trichlorphenylester ist.
 - 26. Verfahren gemäß mindestens einem der Ansprüche 22-25, bei dem der Aminkern von Echinocandin B erhalten wird durch die N-Deacylierung eines natürlich auftretenden cyclischen Hexapeptids.
 - 27. Verfahren gemäß Anspruch 26, bei dem das natürlich auftretende cyclische Hexapeptid Echinocandin B, Tetrahydroechinocandin B, Mulundocandin, L-671 329, S 31794/FI, Sporiofungin oder FR 901379 ist.

45 Revendications

1. Composé de formule (1):

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dans laquelle

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R' est hydrogène, méthyle, ou NH₂C(O)CH₂-;

R" et R" sont indépendamment méthyle ou hydrogène ;

R et R^y sont indépendamment hydroxy ou hydrogène ;

R₁ est hydroxy, hydrogène ou hydroxysulfonyloxy;

R₇ est hydroxy, hydrogène, hydroxysulfonyloxy ou phosphonooxy; et

I) R₂ est un groupe benzoyle substitué représenté par la formule

dans laquelle R₃ est quinolyle ; ou

II) R₂ est un groupe acyle représenté par la formule

dans laquelle

Z est -C≡C-, -CH=CH- ou une liaison carbone-carbone;

A) R₄ est un groupe cycloalkyle en C₃ à C₁₂, bicycloalkyle en C₇ à C₁₀, tricycloalkyle en C₇ à C₁₄, cycloalcoxy en C₃ à C₁₂, naphtyle, pyridyle, thiényle, benzothiényle, quinolyle ou phényle; ou B) R₄ est un groupe phényle substitué par un groupe amino, alkylthio en C₁ à C₁₂, halogène, alkyle en C₁ à C₁₂, alcényle en C₂ à C₁₂, alcynyle en C₂ à C₁₂, alkyle en C₁ à C₁₂ substitué, alcényle en C₂ à C₁₂ substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, phényle substitué, phényle substitué, alcoxy en C₁ à C₁₂, trifluorométhyle, alcoxy en C₁ à C₁₂, trifluorométhyle, alcoxy en C₁ à C₁₂, trifluorométhyle, alcoxy en C₁ à C₁₂, alcoxy en C₁ à C₁₂, trifluorométhyle, alcoxy en C₁ à C₁₂, alcoxy en

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(alkyle en C_1 à C_{12})$

dans laquelle m et n sont des entiers de 2 à 4, et p est 0 ou 1 ; ou

- C) R₄ est un groupe phényle substitué par un alcoxy en C₁ à C₆ substitué par fluoro, bromo, chloro ou iodo; ou
- D) R₄ est un groupe représenté par la formule

-Y-R₆

dans laquelle

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Y est -C≡C- ou -CH=CH-; et

 R_6 est un alkyle en C_1 à C_{12} , alkyle substitué en C_1 à C_{12} ; cycloalkyle en C_3 à C_{12} , bicycloalkyle en C_7 à C_{10} , tricycloalkyle en C_7 à C_{14} , phényle, cycloalcényle en C_3 à C_{12} , naphtyle, benzothiazolyle, thiényle, phényle substitué par amino, alkylthio en C_1 à C_{12} , halogène, alcényle en C_2 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, -O-(C_1) -W-R₅, dans lequel p' est un entier de 2 à 4; W est pyrrolidino, pipéridino ou pipérazino, et R_5 est un hydrogène, alkyle en C_1 à C_{12} , cycloalkyle en C_3 à C_{12} , benzyle ou cycloalkylméthyle en C_3 à C_{12} ; ou alcoxy en C_1 à C_6 substitué par fluoro, bromo, iodo ou chloro; ou

R₆ est un phényle substitué par un groupe polyoxa-alkyle représenté par la formule

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(alkyle en C_1 à C_{12})$$

dans laquelle m, n et p sont tels que définis ci-dessus ; ou

III) R₂ est un groupe acyle représenté par la formule

dans laquelle

Z est -C=C- ou -CH=CH-;

A) R_4 est un hydrogène, alcynyle en C_2 à C_{12} , alcynyle substitué en C_2 à C_{12} , alcoxy en C_2 à C_{12} ; ou B) R_4 est alcoxy en C_1 à C_{12} substitué avec cycloalkyle en C_3 à C_{12} , bicycloalkyle en C_7 à C_{10} , tricycloalkyle en C_7 à C_{14} , alcynyle en, amino, alkylamino en C_1 à C_4 , di-(alkyle en C_1 à C_4)amino, alcanoylamino en C_1 à C_{12} , phényle substitué par un groupe polyoxa-alkyle représenté par la formule

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(alkyle en C_1 à C_{12})$$

dans laquelle m, n et p sont tels que définis ci-dessus ; ou C) R₄ est un alcoxy substitué avec un groupe de formule



dans laquelle R_8 est un alcoxy en C_1 à C_5 éventuellement substitué par un phényle ; ou D) R_4 est un groupe représenté par la formule

-O-(CH₂)_p.-W-R₅

dans laquelle p', W et .R₅ sont tels que définis ; ou

IV) R₂ est un groupe ayant la formule

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dans laquelle Y et R_6 sont tels que définis ci-dessus ; ou V) R_2 est un naphtoyle substitué par R_4 dans lequel

A) R₄ est un cycloalkyle en C₃ à C₁₂, bicycloalkyle en C₇ à C₁₀, tricycloalkyle en C₇ à C₁₄, cycloalcoxy en C₃ à C₁₂, naphtyle, pyridyle, thiényle, benzothiényle, quinolyle ou phényle : ou

B) R_4 est un phényle substitué par un amino, alkylthio en C_1 à C_{12} , halogène, alkyle en C_1 à C_{12} , alcényle en C_2 à C_{12} , alcynyle en C_2 à C_{12} , alkyle substitué en C_1 à C_{12} , alcényle substitué en C_2 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, phényle, phényle substitué, phényle substitué avec un groupe polyoxa-alkyle représenté par la formule

dans laquelle m, n et p sont tels que définis ; ou

C) R_4 est un phényle substitué par un alcoxy en C_1 à C_6 substitué par un fluoro, bromo, chlore ou iodo ; ou D) R_4 est un groupe représenté par la formule dans laquelle

Y a les significations identiques telles que définies ci-dessus ; et

 R_6 est un alkyle en C_1 à C_{12} , alkyle substitué en C_1 à C_{12} ; cycloalkyle en C_3 à C_{12} , bicycloalkyle en C_7 à C_{10} , tricycloalkyle en C_7 à C_{14} , phényle, cycloalcényle en C_3 à C_{12} , naphtyle, benzothiazolyle, thiényle, phényle substitué par amino, alkylthio en C_1 à C_{12} , halogène, alcényle en C_2 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, -O-(CH_2)_{p'}-W-R₅, ou alcoxy en C_1 à C_6 substitué

par fluoro, bromo, iodo ou chloro ; ou R₆ est un phényle substitué par un groupe polyoxa-alkyle représenté par la formule

-O-(CH₂)_m-[O-(CH₂)_n]_p-O-(alkyle en C₁ à C₁₂)

dans laquelle m, n et p sont tels que définis; et leurs sels non toxiques pharmaceutiquement acceptables.

- 2. Composé selon la revendication 1 dans lequel R₁ est hydroxy ou hydrogène et R₇ est hydroxy ou hydrogène.
 - 3. Composé selon la revendication 1 ou la revendication 2 dans lequel R', R" et R" sont un méthyle, R₁ est hydrogène, et R₇ et R^y sont OH.
- 4. Composé selon l'une quelconque des revendications 1 à 3 dans lequel R₂ est de formule

dans laquelle

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Z est une liaison carbone-carbone; et

R₄ est un cycloalcoxy en C₃ à C₇; ou

R₄ est un phényle substitué par un alcoxy en C₁ à C₁₂ ou phényle substitué par un groupe polyoxa-alkyle de formule

-O-(CH₂)_m-[O-(CH₂)_n]_p-O-(alkyle en C₁ à C₁₂);

ΩU

 R_4 est groupe de formule -Y- R_6 , dans laquelle Y est -C \equiv C- ou -CH \equiv CH- et R_6 est un alkyle en C_1 à C_6 , phényle ou phényle substitué par un groupe polyoxa-alkyle de formule

5. Composé selon l'une quelconque des revendications 1 à 3 dans lequel R₂ est de formule

 $- (0) - c - \left(\right) - R$

dans laquelle

50 Z est -C≡C-; et

R₄ est un phényle substitué par un alcoxy en C₁ à C₁₂ ou phényle substitué par un groupe polyoxa-alkyle de formule

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(alkyle en C_1 à C_{12})$$

ou

R₄ est un groupe de formule

-O-(CH₂)_{p'}-W-R₅

dans laquelle W est un groupe pipéridine.

- 6. Composé selon l'une quelconque des revendications 1 à 3 dans lequel R est un hydrogène.
- 7. Composé selon l'une quelconque des revendications 1 à 3 dans lequel R₂ est 4-[4-(phényléthynyl)phényl]benzoyle ou 4-[4-(n-butyléthynyl)phényl]benzoyle.
- 8. Composé de formule (1):

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R" HN RY OH OH OH RY

dans laquelle

R' est hydrogène, méthyle, ou NH₂C(O)CH₂-;
R" et R" sont indépendamment méthyle ou hydrogène;
R et R^y sont indépendamment hydroxy ou hydrogène;
R₁ est hydroxy, hydrogène ou hydroxysulfonyloxy;
R₇ est hydroxy, hydrogène, hydroxysulfonyloxy ou phosphonooxy; et

I) R₂ est un groupe de formule

$$H_3C(CH_2)_2O$$
 $(H_3C)_3CO(CH_2)_2O$
 $H_3C(CH_2)_3O(CH_2)_2O$
 $H_3C(CH_2)_3O(CH_2)_2O$
 $H_3C(CH_2)_3O(CH_2)_2O$
 $H_3C(CH_2)_3O(CH_2)_2O$
 $H_3C(CH_2)_3O$
 $H_3C(CH_2)_3O$
 $H_3C(CH_2)_3O$
 $H_3C(CH_2)_3O$
 $H_3C(CH_2)_3O$
 $H_3C(CH_2)_3O$
 $H_3C(CH_2)_3O(CH_2)_2O$
 et leurs sels pharmaceutiquement acceptables.

- 9. Composé selon la revendication 8 dans lequel R', R" et R" sont un méthyle, R₁ est un hydrogène et R₇ et R^y sont hydroxy.
 - 10. Composé de formule (1):

dans laquelle

R' est hydrogène, méthyle, ou NH2C(O)CH2-;

R" est méthyle ou hydrogène;

R est hydroxy ou hydrogèn ;

R₁ est hydroxy, hydrogène ou hydroxysulfonyloxy

R₇ est hydroxy, hydrogène, hydroxysulfonyloxy ou phosphonooxy;

R₂ est un groupe acyle représenté par la formule

dans laquelle

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Z est -C=C-, -CH=CH- ou une liaison carbone-carbone;

 R_4 est un cycloalkyle en C_3 à C_{12} , bicycloalkyle en C_7 à C_{10} , tricycloalkyle en C_7 à C_{14} , phényle, phényle substitué par amino, alkylthio en C_1 à C_{12} , halogène, alkyle en C_1 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, phényle ou alcoxy en C_1 à C_6 substitué par fluoro, bromo, chloro ou iodo;

ou R₄ est un groupe cycloalcoxy en C₃ à C₁₂;

ou R_4 est un groupe représenté par la formule -Y- R_6 dans laquelle Y est -C \equiv C- ou -CH \equiv CH- et R_6 est un alkyle en C_1 à C_{12} , alkyle en C_1 à C_{12} substitué par phényle; cycloalkyle en C_3 à C_{12} , phényle, cycloalcényle en C_3 à C_{12} , naphtyle, benzothiazol-2-yle ou phényle substitué par amino, alkylthio en C_1 à C_{12} , halogène, alcényle en C_2 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, -O-(CH $_2$) $_p$ -W- R_5 dans laquelle p' est un entier de 2 à 4; W est pyrrolidino, pipéridino ou pipérazino, et R_5 est un hydrogène, alkyle en C_1 à C_{12} , cycloalkyle en C_3 à C_{12} , benzyle ou cycloalkylméthyle en C_3 à C_{12} ; ou alcoxy en C_1 à C_6 substitué par fluoro, bromo, iodo ou chloro ;

ou R2 est un groupe acyle représenté par la formule

dans laquelle Z est -C≡C- ou -CH=CH-;

R₄ est un hydrogène;

 R_4 est un alcoxy en C_1 à C_{12} , alcoxy substitué en C_1 à C_{12} par cycloalkyle en C_3 à C_{12} , bicycloalkyle en C_7 à C_{10} , tricycloalkyle en C_7 à C_{14} , amino, alkylamino en C_1 à C_4 , di-(alkyle en C_1 à C_4)amino, alcanoylamino en C_1 à C_{12} ou un groupe de formule

dans laquelle R_8 est un alcoxy en C_1 à C_6 éventuellement substitué par phényle ; ou R_4 est un groupe représenté par la formule

ou

R₂ est un groupe choisi parmi

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dans laquelle

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Y et R₆ sont tels que définis ci-dessus ; ou R₂ est un naphtoyle substitué par R₄ dans lequel

 R_4 est un cycloalkyle en C_3 à C_{12} , bicycloalkyle en C_7 à C_{10} , tricycloalkyle en C_7 à C_{14} , phényle, phényle substitué par amino, alkylthio en C_1 à C_{12} , halogène, alkyle en C_1 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, phényle ou alcoxy en C_1 à C_6 substitué par fluoro, bromo, chloro ou iodo ; ou R_4 est un groupe représenté par la formule -Y- R_6 dans laquelle Y a les mêmes significations telles que définies ci-dessus et R_6 est un alkyle en C_1 à C_{12} , alkyle en C_1 à C_{12} substitué par phényle; cycloalkyle en C_3 à C_{12} , phényle, cycloalcényle en C_3 à C_{12} , naphtyle, benzothiazol-2-yle, ou phényle substitué par amino, alkylthio en C_1 à C_{12} , halogène, alkyle en C_1 à C_{12} , alcényle en C_2 à C_{12} , alcoxy en C_1 à C_{12} , trifluorométhyle, -O-(CH_2) $_p$ -W- R_5 , ou alcoxy en C_1 à C_6 substitué par fluoro, bromo, iodo ou chloro ;

et leurs sels non toxiques pharmaceutiquement acceptables.

- 11. Composé selon la revendication 10 dans lequel R₁ n'est pas un hydroxysulfonyloxy et R₇ n'est ni un hydroxysulfonyloxy ni un phosphonooxy.
 - 12. Composé de formule

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13. Composé de formule

50 14. Composé de formule

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dans laquelle R est -O(CH₂)₃CH₃, -O(CH₂)₄CH₃, -O(CH₂)₅CH₃, -O(CH₂)₂O(CH₂)₃CH₃ ou -O-(CH₂)₂OC(CH₃)₃.

- 15. Composé selon la revendication 14 dans lequel R est -O(CH₂)₄CH₃.
- 16. Composé selon l'une quelconque des revendications 1-15 à utiliser pour inhiber une activité parasitique.
- 17. Composé selon l'une quelconque des revendications 1-15 à utiliser pour inhiber une activité fongique.
- 18. Composé selon l'une quelconque des revendications 1-15 à utiliser pour inhiber la croissance d'organismes responsables d'infections opportunistes chez les individus immunodéprimés.
 - 19. Composé selon l'une quelconque des revendications 1-15 à utiliser pour inhiber la croissance de <u>Pneumocystis</u> carinii.
 - 20. Formulation pharmaceutique comprenant un composé selon l'une quelconque des revendications 1-15 et un support pharmaceutique approprié.
 - 21. Procédé pour la préparation d'un composé de formule (1) :

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dans laquelle

R' est hydrogène, méthyle, ou NH₂C(O)CH₂-;

R" et R" sont méthyle ou hydrogène;

R est hydrogène;

Ry est hydroxy ou hydrogène;

R₁ est hydroxy ou hydrogène;

R₇ est hydroxy ou hydrogène; et

R₂ est hydrogène ou acyle;

comprenant l'étape consistant à exposer un composé de formule (1) dans laquelle R = OH, à un acide fort en présence d'un agent réducteur, dans un solvant approprié.

22. Procédé de production d'un N-acylhexapeptide cyclique lequel procédé comprend l'acylation d'un noyau amino d'Echinocandin B avec un ester actif d'un acide carboxylique représenté par la formule

HO ______

 $dans \ laquelle \ R \ est \ -O(CH_2)_3CH_3, \ -O(CH_2)_4CH_3, \ -O(CH_2)_5CH_3, \ -O(CH_2)_2O(CH_2)_3CH_3 \ ou \ -O-(CH_2)_2OC(CH_3)_3.$

23. Procédé pour la préparation d'un composé de formule

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dans laquelle R est $-O(CH_2)_3CH_3$, $-O(CH_2)_4CH_3$, $-O(CH_2)_5CH_3$, $-O(CH_2)_2O(CH_2)_3CH_3$ ou $-O-(CH_2)_2OC(CH_3)_3$. lequel procédé comprend l'acylation d'un noyau amino d'Echinocandin B avec un ester actif d'un acide carboxylique représenté par la formule

24. Procédé selon la revendication 22 ou la revendication 23 dans lequel R est -O(CH₂)₄CH₃.

- 25. Procédé selon l'une quelconque des revendications 22-24 dans lequel l'ester actif est un ester de 2,4,5-trichlorophényle.
- 26. Procédé selon l'une quelconque des revendications 22-25 dans lequel le noyau amine d'Echinocandin B est obtenu par N-désacylation d'un hexapeptide cyclique naturel.
- 27. Procédé selon la revendication 26 dans lequel l'hexapeptide cyclique naturel est échinocandin B, tétraéchinocandin B, mulundocandin, L-671 329, S 31794/FI, sporiofungin ou FR 901379.



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- (54) Cyclic peptide antifungal agents and process for preparation thereof.
- 67) Provided are compounds of the formula (1):

$$H_3C$$
 H_3C
 wherein R' is hydrog n, methyl or NH₂C(O)CH₂;

R" is methyl or hydrogen;

R is hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrog n, hydroxysulfonyloxy or phosphonooxy;

R₂ is a novel acyl sid chain. Also provided are nov I formulations, methods of inhibiting fungal and parasitic activity, and a process for preparing did oxy (R=H) forms of the compounds.

Backgr und of th Inv ntion

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This invention relates to cyclic p ptide antifungal agents. In particular, it relates to acyl derivatives of the echinocandin class of cyclic peptide antifungal agents; to methods for treating antifungal and parasitic infections, and to formulations useful in the methods.

The compounds provided by this invention are semi-synthetic antifungal agents in that they are derived from the cyclic peptide antifungals which are produced by culturing various microorganisms. A number of cyclic peptide antifungals are known. Among these are echinocandin B (A30912A), aculeacin, mulundocandin, sporiofungin, L-671,329, FR901379, and S31794/F1. All such antifungals are structurally characterized by a cyclic hexapeptide core, or nucleus, the amino group of one of the cyclic amino acids bearing a fatty acid acyl group forming a side chain off the core or nucleus. For example, echinocandin B has a linoleoyl side chain while aculeacin has a palmitoyl side chain. These fatty acid side chains of the cyclic hexa- peptides can be removed by enzymatic deacylation to provide the free nucleus. (Formula (1), hereinafter, wherein R₂ is hydrogen.) Reacylation of the amino group of the nucleus provides semisynthetic antifungal compounds. For example, the echinocandin B nucleus provides a number of antifungal agents when reacylated with certain unnatural side chain moieties (see *Debono*, U.S. Pat. No. 4,293,489). Among such antifungal compounds is cilofungin which is represented by the formula (1) wherein R is methyl, R₁ is hydrogen and R₂ is p-(n-octyloxy)benzoyl.

Enzymatic deacylation of the cyclic hexapeptides is carried out with deacylase produced by the organism Actinoplanes utahensis and related microorganisms as described by *Abbott et al.*, U.S. Pat. No. 4,293,482.

The present invention provides acylated cyclic hexapeptides having unique side chain acyl groups which, inter alia impart enhanced antifungal and antiparasitic potency e.g. against pathogenic strains of Candida albicans. Also provided is a process for removing the aminal and benzylic hydroxy groups to result in a dideoxy compound of formula (1) (R = H).

Summary of the Invention

The compounds provided by this invention are represented by the following formula (1):

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$$R''' \longrightarrow R' \longrightarrow H$$

$$N \longrightarrow R_2$$

$$N \longrightarrow R'' \longrightarrow N$$

$$N \longrightarrow N $

wherein

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R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" and R" are independently m thyl or hydrogen;

R and RY are indep nd ntly hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy; and

I) R₂ is a substituted benzoyl group repres nted by the formula

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wherein

A) R₃ is a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m and n are integers of from 2 to 4, and p is 0 or 1; or

B) R₃ is an unsaturated hydrocarbon group represented by the formula

wherein Y is -C≡C- or -CH=CH-; or

C) R_3 is a group of the formula -O-(CH₂)_m-G, wherein m is as defined and G is C_7 - C_{10} bicycloalkyl or C_7 - C_{14} tricycloalkyl; or

D) R₃ is quinolyl; or

II) R2 is an acyl group represented by the formula

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wherein

Z is -O-, -C≡C-, -CH=CH-, -CH₂-, -CH₂-, or a carbon to carbon bond;

A) R_4 is hydrogen, C_2 - C_{12} alkynyl, C_2 - C_{12} substituted alkynyl, C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_{14} tricycloalkyl, C_1 - C_{12} alkoxy, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, quinolyl or phenyl; or

B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkenyl, C_1 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkenyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, phenyl, substituted phenyl, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or

C) R₄ is phenyl substituted with C₁-C₆ alkoxy substituted by fluoro, bromo, chloro or iodo; or

D) R_4 is C_1 - C_{12} alkoxy substituted with C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_2 - C_{12} alkynyl, amino, C_1 - C_4 alkylamino, di-(C_1 - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-(O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or

E) R₄ is C₁-C₁₂ alkoxy substituted with a group of the formula



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wherein R₈ is C₁-C₆ alkoxy optionally substituted with phenyl; or

F) R₄ is a group represented by the formula

wher in p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or pip razino, and R_5 is hydrogen, C_1 - C_{12} alkyl, C_3 - C_{12} cycloalkyl, benzyl or C_3 - C_{12} cycloalkylmethyl; or

G) R₄ is a group represented by the formula

wherein Y has the same meanings defined above; and

 R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} substitut d alkyl; C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzothiazolyl, thienyl, indanyl, fluorenyl, phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(CH₂)p'-W-R₅, or C_1 - C_6 alkoxy substituted by fluoro, bromo, iodo or chloro; or

 R_6 is a phenyl substituted by a polyoxa-alkyl group represented by the formula $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein m,n and p are as defined above; or III) R_2 is a group having the formula

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wherein R^x is C₁-C₁₂ alkoxy or a polyoxa-alkyl group represented by the formula $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12}$ alkyl)

wherein m,n and p are as defined above; or IV) R_2 is a group having the formula

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wherein R₉ is phenyl, C₁-C₁₂ alkyl, or C₁-C₁₂ alkoxy; or

V) R₂ is naphthoyl substituted with R₄; and the pharmaceutically acceptable non-toxic salts thereof; with the proviso that when

R' is methyl or NH₂C(O)CH₂-;

R" is methyl;

R" is methyl;

RY is hydroxy;

R is hydroxy; and

either a) or b):

a) R_1 is hydroxysulfonyloxy and R_7 is hydroxy, hydroxysulfonyloxy or phosphonooxy;

b) R₁ is hydrogen or hydroxysulfonyloxy and R₇ is hydroxysulfonyloxy or phosphonooxy;

R₂ is not

i)

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 R_3

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wherein R₃ is

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein p=O; nor

ii)

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wherein Z is a carbon to carbon bond or -O- and R4 is C1-C12 alkoxy; nor

iii) naphthoyl substituted by R₄ wherein R₄ is hydrogen, phenyl, or C₁-C₁₂ alkoxy.

Also provided are formulations and methods for inhibiting parasitic and fungal activity which employ the compounds of the invention, and a process for preparing the dideoxy form of the compounds.

Detailed Description

The term: "C₁-C₁₂ alkyl" r f rs to the straight or branch d chain alkyl hydrocarbon groups such as, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl, t-butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, und cyl and dodecyl groups; and the like.

The term "C₂-C₁₂ alkenyl" refers to groups such as vinyl, 1-propene-2-yl, 1-butene-4-yl, 1-pentene-5-yl, 1-butene-1-yl, and the like.

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The t rm "C2-C12 alkynyl" refers to such groups as ethynyl, propynyl, pentynyl, butynyl and the like.

The term "C1-C12 alkylthio" refers to such groups as methylthio, ethylthio, t-butylthio, and the lik.

The term "C₁-C₁₂ alkoxy" refers to the straight or branched chain oxyalkyl groups such as, e.g. meth xy, ethoxy, propoxy, butoxy, heptoxy, octyloxy, dodecyloxy, and the like.

The term C₃-C₁₂ cycloalkoxy" refers to such groups as cyclopropoxy, cyclobutoxy and the like.

The term "C₃-C₁₂ cycloalkenyl" refers to such groups as cyclopropenyl, cyclobutenyl, cyclopentenyl, and the like.

The term "C₁-C₁₂ substituted alkyl," "C₂-C₁₂ substituted alkenyl", and "C₂-C₁₂ substituted alkynyl", denotes the above substituted one or two times with halogen, hydroxy, protected hydroxy, amino, protected amino, C₁-C₇ acyloxy, nitro, carboxy, protected carboxy, carbamoyl, carbamoyloxy, cyano, methylsulfonylamino, phenyl, substituted phenyl, or C₁-C₁₂ alkoxy.

The term "substituted phenyl" is represented by a phenyl group substituted with one, two, or three moieties chosen from halogen, hydroxy, protected hydroxy, cyano, nitro, C₁-C₁₂ alkyl, C₁-C₁₂ alkoxy, carboxy, protected carboxy, carboxymethyl, hydroxymethoyl, amino, aminomethyl trifluoromethyl or N-(methylsulfonylamino)

The term "C₃-C₁₂ cycloalkyl" refers to such groups as cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl and cycloheptyl.

The term "C₁-C₄ alkylamino" refers to such groups as methylamino, ethylamino, n-butylamino and the like. The term "di-(C₁-C₄ alkyl)amino" refers to such groups as dimethylamino, diethylamino, di-n-propylamino, di-n-butylamino, methyl-n-butylamino, and like tertiary amino groups.

The term ${}^{\circ}C_{1}$ - ${}^{\circ}C_{12}$ alkanoylamino ${}^{\circ}$ refers to such groups as acylamino groups derived from the ${}^{\circ}C_{12}$ carboxylic acids and are exemplified by formamido, acetylamino, propionylamino, butyrylamino, and the like.

The term "C₃-C₁₂ cycloalkylmethyl" refers to those C₃-C₇ cycloalkyls described above further substituted by methyl.

The terms "C₇-C₁₀ bicycloalkyl" and "C₇-C₁₄ tricycloalkyl" refer to such groups as bicyclo[2.2.1.]hept-2-yl, bicyclo[2.2.1.]hep-4-en-2-yl, bicyclo[3.3.1.]nona-3-yl, bicyclo[3.3.1.]nona-2-yl, bicyclo[3.2.1.]oct-2-yl, bicyclo[2.2.2]oct-5-en-2-yl, adamantyl and the like.

The term "dideoxy" refers to compounds of the formula (1) wherein R=H.

The term "inhibiting", such as used in relation to the methods for inhibiting parasitic and fungal activity, is defined to mean its normal definition, i.e., to stop, retard or prophylactically hinder or prevent.

The term "activity", as used in relation to parasitic and fungal activity, includes growth thereof and attending characteristics and results from the existence of the parasite or fungus.

The term "contacting", as used in relation to the methods for inhibiting parasitic and fungal activity by contacting a compound of the invention with a parasite or fungus, is defined to mean its normal definition. However, the term does not imply any further limitations to the process, such as by mechanism of inhibition, and the methods are defined to encompass the spirit of the invention, which is to inhibit parasitic and fungal activity by the action of the compounds and their inherent anti-parasitic and anti-fungal properties, or in other words, the compounds, used in the method are the causative agent for such inhibition.

Examples of acyl groups represented by R_2 in formula (1) are benzoyl substituted by polyoxa-alkyl groups such as, e.g., 2-methoxyethoxy (p=0, m=1), 2-ethoxyethoxy, 2-(2-ethoxyethoxy)ethoxy (m=2, p=1, n=2), 3-(2-ethoxyethoxy)-propoxy, 3-(2-methoxyethoxy)butoxy, and like groups.

Examples of R_3 groups wherein R_2 is benzoyl substituted by an unsaturated hydrocarbon groups -Y-(C_1 - C_{12} -alkyl) include e.g., acetylenic groups -C=C-(C_1 - C_{12} alkyl) and -CH₂=CH₂-(C_1 - C_{12} alkyl) which may be <u>cistors</u> or <u>trans-</u> e.g. propenyl, butenyl, hexenyl, decenyl, and the like; propynyl, butynyl, hexynyl, undecynyl, and like alkynes.

Examples of acyl groups wherein R2 is a group represented by the formula

$$- (0) - C - \sum_{z \in \mathbb{Z}} - Z - \sum_{z \in \mathbb{Z}} - R_z$$

are diphenyl ethers (Z=-0-), diphenyl acetylenes (Z=-C=C-), stilbenes (Z=-CH=CH-), and biphenyls (Z = a carbon to carbon bond). Among xamples of such biphenyl groups, wherein Z is a carbon to carbon bond i.e. a phenyl to phenyl bond, are 4-[4-(butyloxy)phenyl]benzoyl, 4-[4-(cyclobutylm thoxy)-phenyl]benzoyl, 4-[4-cyclopentylmethoxy)phenyl]benzoyl, 4-[4-(cyclohexylethoxy)phenyl]benzoyl, 4-[4-(n-hexyloxy)-phenyl]benzoyl, 4-phenylbenzoyl, 4-[4-(11-amino-und cyloxy)-phenyl]benzoyl, 4-[4-(11-formamidoundecyloxy)phenyl]benzoyl, 4-[4-(isopentyloxy)phenyl]benzoyl, and the like. Examples of such diphenyl ether acyl groups R_2 of the formula abov wherein Z is an oxygen atom are 4-(4-butyloxyphenoxy)benzoyl, 4-(4-hexyloxyphenoxy)benzoyl, 4-(4-hexyloxyphenox

zoyl, 4-(4- thoxyphenoxy)benzoyl, 4-(4-benzyloxyphenoxy)benzoyl, 4-[4-(3-chlorobutyloxy)phenoxy]-benzoyl, 4-(4-dodecyloxyphenoxy)benzoyl, 4-[4-(3-dimethylaminopropoxy)phenoxy]benzoyl and the like. Examples of diph nylacetylene and stilbene acyl groups, R2, wherein Z is an acetylenic bond or an ethylene bond are 4-styrylbenzoyl, 4-(4-methoxystyryl)benzoyl, 4-(4-butyloxystyryl)benzoyl, 4-(phenylethynyl)benzoyl, 4-(4-ethoxyphenylethynyl)benzoyl, 4-(4-cyclohexyloxyphenylethynyl)benzoyl, and the like. Examples of R2 acyl groups represented by the foregoing formula wherein Z is a carbon to carbon bond and R4 is represented by the formula -O-(CH₂)_p-W-R₅ are 4-[4-[2-(N-cyclohexylpiperidine-4-yl)ethoxy]phenyl]benzoyl, 4-[4-[2-(N-hexylpiperidine-4- yl)ethoxy]phenyl]benzoyl, 4-[4-[2-(4-benzylpiperidino)ethoxy]phenyl]benzoyl, 4-[4-[2-(4-cyclohexylpiperidino)- ethoxy]phenyl]benzoyl and like diphenyl acyl groups. Examples of such acyl groups wherein R₄ is represented by the formula -Y-R₆ include 4-[4-(phenylethynyl)phenyl]benzoyl, 4-[4-(phenylethynyl)phenoxy]benzoyl, 4-[4-(hexynyl)phenyl]benzoyl, 4-[4-(styryl)phenoxy]benzoyl, 4-[4-(4-benzylphenylethynyl)-phenyl] benzoyl, 4-[4-[4-4-methylpiperidino)ethoxy]phenylethynyl]phenyl]benzoyl and like acyl groups. Such acyl groups wherein R₄ is represented by the formula -O-(CH₂)_p-W-R₅ form salts of the basic amino groups of the piperidine and piperazine heterocyclic groups with both organic and inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid and phosphoric acid and with organic acids such as the sulfonic acids, benzenesulfonic acid, toluenesulfonic acid, methanesulfonic acid, acetic acid, chloroacetic acid, trifluoroacetic acid, benzoic acid, isophthalic acid, salicylic acid, citric acid, malic acid, succinic acid, malonic acid and like acids.

The following tables contain further examples of the cyclic peptides represented by the formula (1). Table 1 contains examples of cyclic peptides wherein the acyl group R₂ is of the formula

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Table 1

<u>R2</u>

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$$H_{3}C(CH_{2})_{6}O(CH_{2})_{2}O$$
 $H_{3}C(CH_{2})_{7}O(CH_{2})_{2}O$
 $H_{3}C(CH_{2})_{9}O(CH_{2})_{2}O$
 $H_{3}C(CH_{2})_{9}O(CH_{2})_{2}O$
 $H_{3}C(CH_{2})_{3}$
 $H_{3}C(CH_{2})_{3}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$
 $H_{3}C(CH_{2})_{5}$

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The following Table 2 illustrates the compound of the formula (1) wherein R2 is represented by the formula

Table 2

$$\frac{R_{2}}{H_{3}C(CH_{2})_{3}O} \longrightarrow \frac{O}{H_{3}C(CH_{2})_{3}O} \longrightarrow \frac{O}{H_{3}C(CH_{2})_{2}O} \longrightarrow \frac{O}{H_{3}C(CH_{2})_{4}O(CH_{2})_{2}O} \longrightarrow \frac{O}{H_{3}C(CH_{2})_{4}O} \longrightarrow \frac{O}{H_{3}C(CH_{2})_{4}O} \longrightarrow \frac{O}{H_{3}C(CH_{2})_{2}O} \longrightarrow \frac{O}{H_{3}C$$

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Table 2 continued R2

The following Table 3 illustrates compounds of formula 1 wherein R₂ is of the formula as indicated from Table 2 and R₄ is represented by the formula -O-(CH₂)_p-W-R₅.

Table 3

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The acyl cyclohexapeptides represented by formula (1) exhibit antiparasitic activity, for example, they are especially active against the infectious fungi <u>Candida albicans</u> and <u>Candida parapsilosis</u>. They also exhibit significant activity against <u>Aspergillus fumigatus</u>. They are active both <u>in vitro</u> and <u>in vivo</u> and accordingly are useful in combating systemic fungal infections.

The compounds of the invention also inhibit the growth of certain organisms primarily responsible for opportunistic infections in immunosuppressed individuals. For example, the compounds of the invention inhibit the growth of <u>Pneumocystis carinii</u> the causative organism of pneumocystis pneumonia in AIDS sufferers.

The antifungal activity of the compounds of the invention is determined in vitro in standard agar dilution tests and disc-diffusion tests wherein minimum inhibitory concentrations of the test compounds obtained. Stan-

dard in vivo tests in mice ar used to determine the effective dose of the test compounds in controlling systemic fungal infections.

Tables 4A-E below contain the minimum inhibitory concentrati ns (MIC) in micrograms per milliliter (mcg/ml) for compounds of the invention against <u>Candida albicans</u> and <u>Candida parapsilosis</u>, and for certain compounds, the effective dose, ED₅₀, in mice.

In Tables 4A-E, $R'=CH_3$, $R''=CH_3$, $R''=CH_3$, R''=OH, $R_7=OH$ and $R_1=H$, In Tables 4A-D, R=OH, while in Table E, R=H.

In the Table 4A, R₂ is of the formula

with R₃ being as indicated in the Table 4.

In Table 4B, R₂ is of the formula

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where Z is -O- and R₄ is as indicated.

Table 4C is as Table 4B, except Z is a carbon-carbon bond.

Table 4D indicates compound activities in which R₂ is as defined.

In Table 4E, dideoxy (where R=H) compounds are illustrated with R₂ as indicated.

TABLE 4A

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	MIC (MIC (mcg/ml)		
R ₃	C.alb.	C.parap.		
-O(CH ₂) ₂ -O-(CH ₂) ₂ -O-C ₂ H ₅	>20	40	-	
-O-(CH ₂) ₂ -O-C ₅ H ₁₁	>20	40	-	
-O-(CH ₂) ₂ -OC ₇ H ₁₅	10	40	30.3	
-O-(CH ₂) ₂ -O-C ₈ H ₁₇	2.5	80	4.4	
-O-(CH ₂) ₂ -O-C ₁₀ H ₂₁	0.625	5	9.5	
-C≡C-C ₅ H ₁₁	2.5	29	10.5	
-CH=CH-C ₆ H ₁₃ (trans)	0.312	20	4.4	
-C≡C-C ₈ H ₁₇	0.156	10	-	

TABLE 4B

	МІС	(mcg/ml)	ED ₅₀ (mg/kg)
R ₄	C.alb.	C.parap.	
-O-C₄H ₉	>20	40	-
-O-C ₆ H ₁₃	1.25	>20	22.9

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TABLE 4C

		М	ED ₅₀	
5	R4	(mcg C.alb.	/ml) C.parap.	(mg/ml)
	<u></u>		C.parap.	
	-O-C4H9	0.78	10	0.84
10	-O-CH ₂ -cyclobutyl	0.312	10	2.50
	-O-CH ₂ -cyclopentyl	0.039	2.5	1.20
	-O-C5H ₁₁	0.156	0.625	1.86
· 15	-O-C ₆ H ₁₃	0.039	1.25	1.10
· 15	-O-CH ₂ CH ₂ -cyclohexyl	0.039	20	1.6
	$-O-CH_2-CH(C_2H_5)-C_2H_5$	0.039	2.5	4.6
	$-O-CH_2-CH_2-CH(CH_3)_2$	0.309	5	2.00
20	$-O-CH_2-CH_2-C(CH_3)_3$	0.039	2.5	2.21
	-O-(CH ₂) ₂ -O-C ₅ H ₁₁	1.25	20	0.60
	-C≡C-C4H9	0.039	2.5	1.20
25	-C≡C-C ₆ H ₅	0.039	0.625	0.60
	-C ₆ H ₅	0.078	10	1.3
	$-O-(CH_2)_2-N(CH_3)_2$	>20	>20	-
30				
	-O-(CH ₂) ₂ -N	>20	>20	_
	-O-(CH ₂) ₂ -N	F	. 20	2 0
35		5	>20	3.0
	-O-(CH ₂) ₂ -N	0.312	40	0.64
	-O-(CH ₂) ₂ -N	0.039	5	0.24
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TABLE 4D

5		MIC (mcg/ml)			
	<u> </u>	C.alb.	C. parap.		
10	O II -C-(CH ₂) ₄ -O-	40	>80		
15	-C-(CH ₂) ₅ -O	1.25	. 80		
20	-C-CH-O-	0.0039	2.5		
25	-C-CH-O	5	>80		
30	(CH ₂) ₃ CH ₃	80	>80		
35	(CH ₂) ₅ CH ₃	80	>80		
40	(CH ₂) ₁₁ CH ₃	10	>80		
45	O-CH ₂ CH ₃	>80	>80		

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TABLE 4D continued

		MIC (mcg/ml)		
10	O -C	<u>C.alb.</u>	2. parap.	_
15	O-(CH ₂) ₅ CH ₃	20	>80	
20	-C ——O-(CH ₂) ₇ CH ₃	10	>80	
25	O-(CH ₂) ₉ CH ₃			
30		20	>80	
35	O-(CH ₂) ₂ -N -CH ₂ -	20	>80	
	O-(CH ₂) ₅ CH ₃	0.039	5	
40	-C -	0.078	.0.312	

TABLE 4D continued

MIC (mcg/ml)
C.alb. C. 10 15 0.5 80 20 0.005 0.156 25 30 0.039 0.156 **35** 0.156 20 0.005 0.312 45 0.312

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TABLE 4D continued

5	<u> </u>	MIC (mcg/ml C.alb. C) . parap.
10	O-CH ₂	0.312	>80
15			
20	O-CH ₂	0.078	>20

TABLE 4E

_	TABLE 4E			
5		(mc	IC g/ml)	
	R2	C.alb.	C. parap.	
10	-C-C-C4H9	0.039	5.0	
15	- c	>20	1.25	
20	-C-(CH ₂) ₄ -O-(CH ₂) ₄ -O-(C	0.039	2.5	
		>80	>80	
25	O	1.25	40	
30	-C-(CH ₂) ₈ -O-	0.005	2.5	
	-C-(CH ₂) ₁₀ -O	0.0098	0.625	
35	-C-CH-O			
40	-C-CH-O	80	>80	
45	(CH ₂) ₃ CH ₃	20	>80	
	(CH ₂) ₅ CH ₃	40	>80	

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TABLE 4E continued

5	<u>2</u>	MIC (mcg/ml) C.alb. C. parap.	
10	O II -C-CH-O	2 25	
15	c	1.25 >80	
20	O-CH ₂ CH ₃ O-(CH ₂) ₅ CH ₃	·80 :-80	
25	O -C	10 >80	
30	O -C	10 >80	
35		5.0 >80	
40	O-(CH ₂) ₂ -N -CH ₂ -	1.25 >80	
4 5	O-(CH ₂) ₅ CH ₃	0.078 1.25	
	O-(CH ₂) ₇ CH ₃	0.039 0.125	

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TABLE 4E continued

5		MIC (mcg/ml)			
	<u>R2</u>	C.alb.	C. parap.		
10	-C-(CH ₂) ₉ CH ₃	0.156	0.625		
15	$-C - CH_2 - CH$	0.156	5.0		
20					
25		0.625	80		
30		0.005	0.156		
35					
		0.039	0.156		

The non-dideoxy compounds of the invention (formula (1) are prepared with the amino nuclei of the cyclic hexapeptides which are represented by the formula when R_2 is hydrogen. These amino nuclei are obtained from the known natural products by the known enzymatic deacylation by which the fatty acid side chains of the natural compounds are removed. For example, echinocandin B which can be represented by the formula (1) wherein R'=R''=R'''=methyl, R is OH, R' is hydroxy, R_1 is H, R_7 is OH, and R_2 is linoleoyl, is deacylated to provide the echinocandin B nucleus ($R_2=H$) with the deacylase produced by the organism Actinoplanes utahensis as described by U.S. Patent Nos. 4,293,482 and 4,304,716.

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The known natural cyclic hexapeptides which are N-deacylated to provide the amino nuclei starting materials include echinocandin B (also known as A-30912A), aculeacin (palmitoyl side chain), tetrahydoechinocandin B (stearoyl side chain), mulundocandin (branched C_{15} side chain), L-671,329 (C_{16} branched side chain), S 31794/F1 (tetradecanoyl side chain), sporiofungin (C_{15} branched side chain) and FR901379 (palmitoyl side chain). The amino nuclei obtain d by the N-deacylation are then acylated by employing known amino acylation procedures to provide the N-acyl cyclic hexapeptides represented by the formula (1) wherein R_2 represents the acyl groups defined hereinabov. The acylating moiety is preferably an active ester of the carboxylic acid RCOOH such as the 2,4,5-trichlorophenyl ester. The R_2 COOH precursor acids are prepared by the hydrolysis of the nitrile R_2 CN or the ester R_2 COOC₁-C₄ alk. These nitrile and ester intermediates are prepared by known

methods.

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The alkoxy aromatic (ie. phenyl and biph nyl) compounds of Tabl s 5-10 are prepared by one of the two following procedures:

A. The hydroxyaromatic compound (1 equivalent) is dissolved in acetonitrile (200-300 ml) and a base, such as potassium t-butoxide or potassium carbonate, (1-equivalent), is added. An alkyl bromide, iodide, or p-toluenesulfonate (1 equivalent) is then added and the solution is refluxed for 6 hours. The solvent is evaporated in vacuo and the residue is dissolved in ether and 2N sodium hydroxide. The ether layer is dried over magnesium sulfate and evaporated to give the alkoxyaromatic product.

B. The hydroxyaromatic compound (1 equivalent), alkyl alcohol (1 equivalent), and triphenylphosphine (1 equivalent) are dissolved in tetrahydrofuran (200-300 ml) and diethylazodicarboxylate (1 equivalent) is added dropwise over 10 minutes at room temperature. After 17 hours the solvent is removed in vacuo and the residue is dissolved in ether. This organic layer is extracted with 2N sodium hydroxide solution, dried over magnesium sulfate, and evaporated to give a product which is crystallized from ether/pentane or, if the product contains a tertiary amine, the hydrochloride salt is formed and crystallized from methanol/ethyl acetate.

5 10	BrCH ₂ CH(CH ₂ CH ₃) ₂ I(CH ₂) ₅ CH ₃ Br(CH ₂) ₂	Br(CH ₂) ₄ CH ₃ CH ₃ SO ₃ -CH ₂ CH ₃ SO ₃ -(CH ₂) ₂ C(CH ₃) ₃	I(CH ₂) ₃ CH ₃ CH ₃ SO ₃ -CH ₂ Br(CH ₂) ₂ CH(CH ₃) ₂ CH ₃ SO ₃ -(CH ₂) ₂ O(CH ₂) ₄ CH ₃	Alkyl halide or tosylate
	8.5 10.8 4.2	15.3 13.0	9.4 12.3 7.7 7.6	8 V
20	> > >	> >>	>> >>	Meth
25				hod
30	-CH ₂ CH(CH ₂ CH ₃) ₂ -(CH ₂) ₅ CH ₃ (CH ₂) ₂	-(CH ₂) ₄ CH ₃ CH ₂ (CH ₂) ₂ C(CH ₃) ₃	-(CH ₂) ₃ CH ₃ -(CH ₂) ₃ CH ₃ -(CH ₂) ₂ CH(CH ₃) ₂ (CH ₂) ₂ O(CH ₂) ₄ CH ₃	TABLE 5
35	2CH ₃) ₂	— _{— ,}	3 (CH ₃) ₂) ₄ CH ₃	J _R
40	CN CN CO ₂ CH ₃	X X X X C C C	C N C N C C N C C N C C N C C N C C N C C N C C N C C N C C N C C N C C N C C N C	R ₂
45	3.0 11.4 4.5	20.3 12.2 11.8	3.2 5.3 9.2 4.8	₩t.

5 10	HO(CH ₂) ₂ -N CH ₂ -CH ₂ -C	HO(CH ₂) ₂ -N NCH ₂	$HO(CH_2)_2-N$ CH_2 $HO(CH_2)_2$ $N(CH_2)_5CH_3$	HO(CH ₂) ₂ -N -(CH ₂) ₂ CH ₃	Alcohol	
•	9.3	0.5	0.5	3.6	g Wt.	
	&	B B	B 5	B	Meth	
25					hod	
30	· (CH ₂) _Z -N	(CH ₂) ₂ -N	(CH ₂) ₂ -N	(CH ₂) ₂ -N		TABLE 6
35		Z () (NCH)	CH ₂ CH ₃	CH ₂) ₂ CH ₃	R	
40			Î	₂ СН ₃		8
45	9.6	0.5	0.8	6.2	W.L.	

5 10	I(CH ₂) ₃ CH ₃ I(CH ₂) ₅ CH ₃		HOCH ₂ (CH ₂) ₃ CH ₃	CH ₃ SO ₃ -(CH ₂) ₂ O(CH ₂) ₉ CH ₃	CH ₃ SO ₃ -(CH ₂) ₂ O(CH ₂) ₇ CH ₃	CH3 SO3-(CH2)2O(CH2)6CH3	Tosylate or alcohol	
	6.1 6.1		10.0	27.1	25.8	23.4	wt.	
25	Method A		œ	>	>	>	Method	
30	RO	TABLE 8	CH ₂ -	-(CI	-(CI	-(CI	PO P	TABLE 7
35	-(CH ₂) ₃ CH ₃ -(CH ₂) ₅ CH ₃		(СН ₂) ₃ СН ₃	-(CH ₂) ₂ O(CH ₂) ₉ CH ₃	-(CH ₂) ₂ O(CH ₂) ₇ CH ₃	-(CH ₂) ₂ O(CH ₂) ₆ CH ₃	R OCH ₂ CH ₃	
40	OCH3		¥	13		13	Ï	
45	WL. 12.3 4.7		13.6	21.0	7.9	20.9	wt.	

5 10	H ₃ C \ SO ₃ -(CH ₂) ₂ OC(CH ₃) ₃	$H_3C \left(\right) SO_3 \cdot (CH_2)_2O(CH_2)_3CH_3$	Alkylhalide or tosylate			H ₃ C SO ₃ -(CH ₂) ₂ OC(CH ₃) ₃	H_3C $SO_3 \cdot (CH_2)_2O(CH_2)_3CH_3$	Alkylhalide or tosylate		
20	4.9	3.8 6	7 × 1			2.7	2.6 2.7	g Wt.		
25	\	> >	Method		Table 10	>	> >	Method		Table 9
30	—(СН ₂) ₂ ОС(СН ₃) ₃	-(CH ₂) ₂ CH ₃ -(CH ₂) ₂ O(CH ₂) ₃ CH ₃	R	RO ()		-(сн ₂) ₂ ос(сн ₃) ₃	-(CH ₂) ₂ CH ₃ -(CH ₂) ₂ O(CH ₂) ₃ CH ₃	· IR	RO \\	
35	H ₃) ₃	3 3 3 CH ₃				H ₃) ₃	3 3 CH3			
40	5.2	5.1	Wt.	OCH3		2.6	4.4 2.6	e Kt	OCH ₃	

The alkynyl and alkenyl aromatic compounds contained in Tables 11-14 are prepared by the following procedure:

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An aromatic bromide, iodide, or trifluoromethane-sulfonate (1 equivalent) is dissolved in acetonitrile (600 ml/0.1 mole of aromatic reactant) under a nitrogen atmosphere. An alkyne or alkene (1 equivalent), triethylamine (2 equivalents), palladium dichloride (0.05 equivalents), triphenylphosphine (0.1 equivalents), and cuprous iodide (0.025 equivalents) are added and the solution is refluxed for 17 hours. The solvent is removed in vacuo and the residue is slurried in ether (300 ml). Solids ar removed by filtration and the filtrate is washed with 1N hydrochloric acid solution. The organic layer is dried over magnesium sulfate and evaporated to yill the product.

5	_		
10	Acetylene H——(CH ₂),CH ₃	Acetylene H=-(CH ₂) ₃ CH ₃ H=-Si(CH ₃) ₃	Acetylene or olefin H=-(CH ₂) ₅ CH ₃ H=-(CH ₂) ₇ CH ₃ H=-SI(CH ₃) ₃
15	₩ t .	8 1.8 1.4 1.4	E WL 8 12.1 6.1 15.2 1.9
20	Br Wt.	6.0 40.0	
25	ocH.	E OCH3	J J 3
30		TABLE 13	TABLE 11 -CH: TABLE 12
35	R O O O O O O O O O O O O O O O O O O O	-с=-(CH ₂) ₃ CH ₃ -SI(CH ₃) ₃	C=-(CH ₂) (CH ₂) (CH ₂) (CH ₂) (CH ₂) (CH ₂) (CH ₂)
40		을	ans)
	= 00 =	₩£ 2.6 2.3	WL 8 26.2 0.6 28.1 1.9
45			

$H = \left(\right) \right)^{2}$ och ₃	1.2 Br-//	$H = \sqrt{\frac{2}{3}}$ осн ₃ 22.2 вг $\sqrt{\frac{2}{3}}$ он	$H = \sqrt{\frac{1}{2}} ^{2} OCH_{3}$ 10.5 $I = \sqrt{\frac{1}{2}} OH$	Acetylene wt. Halide	IABLE 14
C=C-{\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	į		9.7 HO \\ ____\\\\\\\\\\\\\\\\\	Product	
	1.5	19.4	10.2	P W	

The aromatic boronic acids listed in Table 15 were prepared by th following procedure:

An aromatic halid (1 equivalent) is cooled to -78°C in t trahydrofuran solvent. Butyl lithium (1.2 equivalents) is added. After 15 min triisopropyl borate (2 equivalents) is added and after 10 min of stirring th cooling bath is remov d. When the reaction has warmed to room temperature water is added to quench the reaction followed by 1N HCI. The organic layer is removed under reduced pressure leaving a solid precipitate which is collected by filtration. This solid is washed with hexane leaving the pure boronic acid.

The terphenyl esters listed in Table 16 were made in the following manner:

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An aromatic boronic acid (1 equivalent), methyl 4-iodobenzoate (1 equivalent), and potassium carbonate (1.5 equivalents) were mixed in a nitrogen-purged toluene solution. Alternatively, the trichloro phenyl ester of iodobenzoate my be used. Added tetrakis(triphenylphosphine)palladium (0.03 equivalents) and refluxed for 7 hrs. The solution was decanted to remove the potassium carbonate and reduced in vacuo. The residue was triturated with acetonitrile and the product solid was collected by filtration.

20	-0(CH ₂) ₃ CH ₃ -0(CH ₂) ₄ CH ₃ -0(CH ₂) ₅ CH ₃ -0(CH ₂) ₂ O(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ OC(CH ₃) ₃	R		R Och	H Oolch	R Oolch	R Och	H O(C)		
25 30	5.0 6.0 3.4 3.7	(HO) ₂ B - W _{1.} (g)		O(CH ₂) ₂ OC(CH ₃) ₃	0(CH ₂) ₂ 0(CH ₂) ₃ CH ₃	O(CH ₂) ₅ CH ₃	O(CH ₂) ₄ CH ₃	O(CH ₂) ₃ CH ₃		
35	3.2 3.7 2.8 3.6	}	TABLE 16	5.0	13.6	10.9	31.0	10.6	R=Br Wt. (g)	TABLE 15
4 5	2 3 5	-I H ₃ co Д			5	4		6.	R=B Wt	
50	4.2 5.2 3.5 3.7 2.2			. 9	.7		2.0		$R=B(OH)_2$ $W_1. (g)$	

The aromatic nitriles or carboxylate esters described in Tabl s 5-16 can be converted to carboxylic acids

by one of the two following hydrolysis procedures:

A. An aromatic nitrile is dissolved in ethanol and an excess of 50% sodium hydroxide solution and r fluxed for 2 hours. Water is added until a solid precipitat s. The precipitate is collected by filtration, added to dioxane and 6N hydrochloric acid solution and refluxed for 17 hours. Water is added and the carboxylic acid product crystallizes and is collected by filtration and dried under vacuum.

B. A carboxylate methyl ester is dissolved in methanol, excess 2N sodium hydroxide solution is added and the solution is refluxed for 5 hours. The solution is made acidic with excess hydrochloric acid and water is added until a precipitate forms. The carboxylic acid is collected by filtration and dried under vacuum.

The carboxylic acids are converted to 2,4,5-trichlorophenyl esters shown in Tables 17-25 by the following general procedure:

The aromatic acid (1 equivalent), 2,4,5-trichlorophenol (1 equivalent), and N,N'-dicyclohexylcarbodiimide (1 equivalent) are dissolved in methylene chloride. The mixture is stirred for 17 hours after which it is filtered. The filtrate is evaporated to dryness and the residue is dissolved in ether, filtered, and pentane is added until crystallization begins. The crystalline product is collected by filtration and dried under vacuum.

. 15

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5 10 15	$-(CH_2)_2 - N \longrightarrow CH_2 \longrightarrow$	-(CH ₂) ₂ - N	$-(CH_2)_2$ $N-CH_2$	-(CH2)2 - N-(CH2)5CH3	-(CH ₂) ₂ -N CH ₂ -	-(CH2)2-N -(CH2)2CH3	·(CH ₂) ₂ -	-(CH ₂) ₂ C(CH ₃) ₃ -CH ₂ CH(CH ₂ CH ₁) ₂ -(CH ₂) ₅ CH ₁	-CH ₂	-(CH2)2CH(CH3)2 -(CH ₂) ₂ O(CH ₂) ₄ CH ₃ -(CH ₂) ₄ CH ₃	-(CH ₂) ₃ CH ₃		HO POH POH POH	
25	7.5	7.2	2.0	1.0	3.0	3.3	ب د در ا	2.3 1.5	4.4	3.0 2.2 5.7	1.9 4.2			
30						•			•				2.4.5-1	TABLE 17
35	7.3	0.8	0.8	1.0	2.3	1.5	1.0	2.6 0.8	<u></u>	1.7 1.3 5.1	4.4	g TM	2.4.5-trichlorophenol	
40 45												7.1 K-1	- 1	

5 10	-(CH ₂) ₂ 0(CH ₂) ₆ CH ₃ -(CH ₂) ₂ 0(CH ₂) ₇ CH ₃ -(CH ₂) ₂ 0(CH ₂) ₉ CH ₃ -CH ₂ (CH ₂) ₃ CH ₃	!R	-C=-(CH ₂) ₃ CH ₃
15		RO OH	
20			모 오
	5.6 7.8 4.0	? ₹	. 1 2.0 ¥1.
25		į.	
30		TABLE 19 2.4	TABLE 18 2.4
35		5-tric	.5-tric
40	8 6.6 1.3	henol	2.4.5-trichlorophenol wt. g 0.6
45		ester	ester

5 10 15	FO 7 1 1	Carboxylic acid	-(CH ₂) ₃ CH ₃ -(CH ₂) ₅ CH ₃	-C=-(CH ₂) ₅ CH ₃ -C =-(CH ₂) ₅ CH ₃ (trans) -(C=-(CH ₂) ₇ CH ₃	R
20	0.8 8.3 8.3		О————————————————————————————————————	4.6 1.2 11.1 1.5	POH WI.
30		TABLE 22 2.4.5-tr	TABLE 21 2.4.5-11		TABLE 20 2.4.5-tr
35 40	B 13.2 1.2	2.4.5-trichlorophenol ester	2.4.5-trichlorophenol ester wt. g 1.4 2.4	3.5 0.5 13.2 1.5	2.4.5-trichlorophenol ester
	1	 	1 14 1		

5	-0(CH ₂) ₂ O(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ OC(CH ₃) ₃	R	-0(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ O(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ OC(CH ₃) ₃	□	-0(CH ₂) ₃ CH ₃ -0(CH ₂) ₄ CH ₃ -0(CH ₂) ₅ CH ₃ -0(CH ₂) ₂ O(CH ₂) ₃ CH ₃ -0(CH ₂) ₂ OC(CH ₃) ₃	₹
15		₩ W				. 공
20	2.9 2.0 2.0	W1. (g)	6 9 5	Wr. (e	3.3 3.0 2.3 1.3	ST-TM
25		TABLE 25		TABLE 24		TABLE 23
30						23
35	2.5 1.5 1.3	2,4,5-Trichlorophenol	5.2 5.2 2.1	2,4,5-Trichlorophenol	4.8 2.5 3.9 4.4 1.9	2,4,5-Trichlorophenol Wt. (g)
40	_ 5. 5.	phenol	2	phenol	8 2 9 4 9	phenol
45		ester		ester		ester

The dideoxy compounds of formula (1) are prepared by removing the benzylic and aminal hydroxy groups. The process includes subjecting a non-dideoxy compound of formula (1) (wherein R_2 may be hydrogen or acyl) to a strong acid such as trichloroacetic acid, trifluoroacetic acid or borontrifluorid—therate with trifluoroacetic acid being preferred, and a reducing agent, such as sodium cyanoborohydrid—or triethylsilane, with triethylsilane being preferred. The reaction takes plac—at temp ratur—s of betwe—n -5 and 70°C, and in a suitable solvent such as m—thylen—chloride, chloroform or ac—tic acid, with dichloromethan—being pr—ferred. Th—acid should be pres—nt in an amount of 2 to 60 moles per mole of substrat—, and the reducing agent should be pr—s—nt in an amount of 2 to 60 moles per mole of substrate. This proc—ss affords selective removal of the aminal and

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benzylic hydroxy groups.

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The compounds represented by the formula (1) have improved properties over the previously known N-acyl hexap ptide antifungals. For xample, in general th compounds exhibit oral bioavailability, a property which is important for any systemic antifungal agent. Also, numerous N-acyl compounds of the formula (1) have enhanced antifungal activity and enhanced water solubility.

Among the N-acyl hexapeptides represented by the formula (1) certain are preferred embodiments of the invention. The compounds wherein R₂ is a diphenyl acyl group

$$- C(O) - Z - Z - R_{2}$$

wherein Z is a carbon to carbon bond and R_4 is an alkoxy, cycloalkoxy or cycloalkylalkoxy group are preferred antifungals. Also preferred compounds are represented when Z is a carbon to carbon bond and R_4 is -Y-R₆ and R₆ is C₁-C₁₂ alkyl phenyl or substituted phenyl and Y is an acetylenic bond.

A further preferred group of N-acyl hexapeptides is represented when Z is a carbon to carbon bond and R_4 is represented by -O-(CH₂)_p-W-R₅ and wherein W is a piperidine group.

Examples of preferred compounds of the above first mentioned group include 4-(4-alkoxyphenyl)benzoyl wherein the alkoxy group is preferably a C_5 - C_{10} alkoxy group or C_1 - C_4 alkoxy substituted by C_3 - C_7 alkyl. Examples of such preferred compounds are represented by the formula 1 wherein R_2 is 4-(4-n-hexyloxyphenyl)benzoyl, 4-(4-n-heptyloxyphenyl)benzoyl, 4-(4-n-octyloxyphenyl)benzoyl, 4-[4-(3,3-dimethylbutoxy)phenyl]benzoyl, 4-[4-(2-cyclopentyl-ethoxy)phenyl]benzoyl and 4-[4-(2-cyclopexyloxyethoxy)phenyl]benzoyl.

Examples of the second above mentioned preferred compounds wherein R_4 is -Y- R_6 include 4-[4-(phenylethynyl)phenyl]benzoyl and 4-[4-(n-butylethynyl)phenyl]benzoyl.

Examples of preferred compounds of the invention wherein R_4 represents -O-(CH₂)_p-W-R₅ are represented when R_2 has the formula

wherein W-R₅ is piperidino, 4-n-propylpiperidino, 4-benzylpiperidino, 4-cyclohexylpiperidino, 4-cyclohexylmethylpiperidino, and the pharmaceutically acceptable acid addition salts such as the hydrochloride salts, the sulfate salts or the phosphate salts.

Preferred cyclohexylpeptide compounds are represented by the formula 1 wherein R'=R"= methyl, R₁ is hydrogen and R₂ is a preferred acyl group as defined hereinabove.

Table 26 is a list of the most preferred R_2 substituents, wherein $R=R_7=R^Y=OH$; $R'=R''=CH_3$; and $R_1=H$.

45	40	<i>30 35</i>	25	15 20	5
					* m+1; ** m+ Li +
				C=C-{\}-{\}-\\	
1166.4758*	0.2	2.6	1.8		
1170.5261*	1.4	2.9	1.9		(H ₃ C) ₃ CO(CH ₂) ₂ O
1170.5234*	6.5	6.7	4.4		H ₃ C(CH ₂) ₃ O(CH ₂) ₂ O
1154.5343*	1.4	. 5.0	3.5		н₃с(сн₂)₅о
1140.5103*	5.1	3.7	2.5		H3C(CH2)40
1126.5025*	1.3	7.4	4.6		н₃с(сн₂)₃о
1194.5247*	2.4	1.5	1.3	10-C2	(H₃C)₃CO(CH₂)₂O
1194.5213*	3.0	3.2	2.0		H₃C(CH₂)₃O(CH₂)₂O ⟨
1136.4832*	0.9	3	2.4		н₃с(сн₂)₂о⟨⟩
1194.5282*	1.1	6.4	5.2		H ₃ C(CH ₂) ₃ O(CH ₂) ₂ O
1200.5336**	2.0	2.5	2.1		(H₃C)₃CO(CH₂)₂O €
1142.4951**	1.4		5.2		н₃с(сн₂)₂о⟨⟩
FABMS	Product (g)	A30912A Nucleus (g)	Ester Reactant (g)	R ₂	
			I VOLT 70		

The N-acylhexap ptid s provided by this invention are useful in the treatment of fungal infections both systemic infections and skin infections. Accordingly this invention also provides a method for treating fungal infections in man and animals which comprises administering to said host an antifungally frective non-toxic amount of an N-acyl-cyclohexapeptide represent d by the formula 1. A preferred antifungal method comprises

administering an N-acylhexapeptide compound where, in formula 1, R'=R"= methyl, R_1 is hydrogen and R_2 is a preferred acyl group as defined her inabove.

The antifungal compound can be administered parenterally, e.g. i.m., i.p. or s.c., nasally, orally or can be applied topically for skin infections. The dose administered of course will vary depending on such factors as the nature and severity of the infection, the age and general health of the host and the tolerance of a particular host to the particular antifungal agent. The particular dose regimen likewise may vary according to such factors and may be given in a single daily dose or in multiple doses during the day. The regimen may last from about 2-3 days up to about 2-3 weeks or longer.

This invention also provides pharmaceutical formulations useful for administering the antifungal compounds of the invention. These formulations comprise an N-acylhexapeptide represented by the formula 1 or a pharmaceutically acceptable, non-toxic salt thereof and a pharmaceutically acceptable carrier.

For parenteral administration the formulation comprises a compound of the formula 1 and a physiologically acceptable diluent such as deionized water, physiological saline, 5% dextrose and other commonly used diluents. The formulation may contain a solubilizing agent such as a polyethylene glycol or polypropylene glycol or other known solubilizing agent. Such formulations may be made up in sterile vials containing the antifungal and excipient in a dry powder or lyophilized powder form. Prior to use, the physiologically acceptable diluent is added and the solution withdrawn via syringe for administration to the patient. For oral administration, the antifungal compound is filled into gelatin capsules or formed into tablets. Such tablets also contain a binding agent, a dispersant or other suitable excipients suitable for preparing a proper size tablet for the dosage and particular antifungal compound of the formula 1. For pediatric or genatric use the antifungal compound may be formulated into a flavored liquid suspension, solution or emulsion. A preferred oral carrier system is lineolic acid, cremophor RH-60 and water and preferably in the amount (by volume) of 8% lineolic acid, 5% cremophor RH-60, and 87% sterile water. The compound is added to the system in an amount of 2.5 to 40 mg/ml.

For topical use the antifungal compound can be formulated with a dry powder for application to the skin surface or it may be formulated in a liquid formulation comprising a solubilizing aqueous liquid or non-aqueous liquid, e.g., an alcohol or glycol. Such formulations are useful forms for use in the antifungal method provided herein.

The N-acylcyclohexapeptides provided herein may be formulated as described above in unit dosage formulations comprising for injection between about 50 mg and about 500 mg per vial. For oral use gelatin capsules or tablets comprising between about 100 mg and about 500 mg per capsule or tablet can be provided.

Preferred formulations of the invention comprises the active ingredient presented by the formula 1 wherein R'=R''= methyl, R_1 is hydrogen and R_2 is 4-[4-(phenylethynyl)-phenyl]benzoyl in gelatin capsules or as active ingredient the antifungal represented by the formula 1 wherein R'=R''= methyl, R_1 is hydrogen and R_2 is 4-[4-[2-(4-cyclohexyl-piperidino)ethoxy]phenyl]benzoyl or the hydrochloride salt form thereof in tablet or gelatin capsules. Further preferred formulations are those in which a preferred compound, as described above, is employed.

In yet a further aspect of the present invention there is provided a method for treating patients suffering from Pneumocystis pneumonia. The method can be used prophylactically to prevent the onset of the infection which is caused by the organism Pneumocystis carinii. The N-acylcyclicpeptide can be administered parenterally, e.g. via intramuscular (i.m), intravenous (iv.) or intraperitoneal (i.p.) injection, or orally or by inhalation directly into the airways of the lungs. Preferably the cyclic peptide is administered via inhalation of an aerosol spray formulation of the compound.

An effective amount of a cyclic peptide will be between about 3 mg/kg of patient body weight to about 100 mg/kg. The amount administered may be in a single daily dose or multiple doses e.g. two, three or four times daily throughout the treatment regimen. The amount of the individual doses, the route of delivery, the frequency of dosing and the term of therapy will vary according to such factors as the intensity and extent of infection, the age and general health of the patient, the response of the patient to therapy and how well the patient tolerates the drug. It is known that PCP infections in AIDS patients are highly refractory owing to the nature of the infection. For example, in severe, advanced infections the lumenal surface of the air passages becomes clogged with infectious matter and extensive parasite development occurs in lung tissue. A patient with an advanced infection will accordingly require higher doses for longer periods of time. In contrast, immune deficient patients who are not severely infected and who are susceptible to PCP can be treated with lower and less frequent prophylactic d s s.

Th activity of th cyclicpeptide represented by the formula 1 is d monstrated in immunosuppressed rats. The tests were carried out in general as follows. One week after initiation of immunosuppression rats were inoculated intratracheally with parasites and maintained on immunosuppression for the remainder of the study. Prophylactic treatments began on day after parasite inoculation and therapeutic treatments began 3 or 4 weeks later after moderat PCP d v loped. Eight or ten animals were assigned to the following groups: thos

receiving test compound; non-tr ated <u>Pneumocystis</u> infected control animals; animals treated with trimetho-prim-sulfamethoxazole (TMP-SMX); or non-treated, non-infected control animals. The efficacy of different treatments was evaluated by monitoring animal weights and survival during the studies and by determining the severity of PCP at necropsy. Stained impression smears of the lungs and stained lung homogenates were evaluated to determine the intensity of P. carinii infection.

The immune deficient rats employed in the tests were prepared as follows. Female Lewis rats weighing from 120-140 g each were immune suppressed with methyl prednisolone acetate at a dose of 4 mg/100 g for the first week, 3 mg/100 g for the second week and continuing weekly thereafter at 2 mg/100 g. All rats, except for the non-infected control rats, were inoculated intratracheally with 0.1 ml to 0.2 ml of Dulbecco's Modified Eagle Media containing between >10⁵ and 10⁶ P. carinii (trophozoites, precysts and cysts) harvested from the lungs of heavily infected donor animals (infection scores of 6) and maintained as cryopreserved (liquid nitrogen) inocula. Rats were maintained on immune suppression and PCP was allowed to develop for 3 or 4 weeks before initiation of therapy with test compounds. Body weights were recorded weekly and rats were allocated into treatment groups such that each group had a similar distribution of percent weight loss among animals. Rats were treated with test compounds for 2 or 3 weeks and then were necropsied. For prophylaxis studies, administration of test compound was initiated one day after intratracheal inoculation of parasites and was continued until the rats were necropsied.

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Following the evaluation period for test compounds, the rats were necropsied and test results evaluated by Giemsa-stained, silver-methenamine stained impression smears and/or by silver-methenamine stained lung homogenate (see below). Necropsy was carried out as follows. The test rats were anesthetized with a mixture of ketamine hydrochloride and xylazine and then exsanguinated via the right atrium. Internal organs in the abdominal and thoracic cavities were examined for gross lesions.

A small portion of lung tissue from the left lobe of each rat was used to make the impression smears described below. Giemsa-stained impression smears were evaluated to determine the total number of parasites (trophozoites, precysts, and cysts). Impression smears from rats in groups whose treatments exhibited some anti-Pneumocystis activity (as judged by infection scores from Giemsa-stained slides) and from rats in the control groups were also stained with methamine silver, a stain specific for the cyst wall of the organism. Impression smears were randomized, numbered, and then evaluated. The infection scores used were as follows:

Score	Basis
0	No parasites found
1	1 to 5 parasites/10 oil fields
2	ca 1 parasite/field
3	2-10 parasites/field
4	>10 but <100 parasites/field
5	>100 but <1,000 parasites/field

A score of 6 was reserved for those infections with impression smears containing >1,000 organisms/field (too numerous to count). Giemsa-stained slides were examined microscopically using a final magnification of 1008X. Methenamine silver-stained slides were examined with a final magnification of 400X.

Cysts in rat lung tissue were quantified as follows. A small portion of lung tissue from the left lobe of each rat was used to make impression smears as described above. The remainder of each lung was weighed, placed in a tube containing Hanks balanced salt solution (HBSS) (40X the lung weight) and homogenized using a Biinkman model tissue homogenizer. Two $\mu1$ samples of the homogenized lung samples (1:4 dilution in HBSS) were placed in wells of teflon-coated, 12-well slides, stained with methenamine silver, and the number of cysts were scored as described above for the impression smears.

The activity and efficacy of two preferred N-acylcyclohexapeptides in the test animals is presented below. The compound of the formula 1 wherein R'=R"= methyl, R₁ is hydrogen and R₂ is 4[(4-phenylethynyl)phenyl]benzoyl when administ red as an aerosol solution at a concentration of 5 mg/ml for one hour, twice we kly for 5 weeks resulted in 90% reduction in <u>P. carinii</u> cysts in the lungs. When given orally at 10 mg/kg, bid for 3 weeks, the number of cysts in the lungs was reduced by >99% when compared with infected vehicle controls.

When the preferred N-acylcyclicpeptides were administered orally and by intraperitoneal injection the compound was ffective in clearing P. carinii cysts from the lungs of heavily infected rats. For example, when the

comp und was administered at 10 or 40 mg/kg, bid for 4, 8 or 12 days, the number of identifiable cysts in the lungs of heavily infected rats was reduced by >99%. Similar efficacy was observed when the compound was administered i.p. at 1 mg/kg.

When tested orally for prophylactic activity, the preferred compound exhibited >99% cyst reduction in one of two studies when infected animals were dosed at 1 mg/kg and when given higher doses of 5 or 4 mg/kg.

Another preferred compound of the invention represented by the formula 1 wherein R'=R"= methyl, R_1 is hydrogen and R_2 is 4-[4-[2-(4-cyclohexylpiperidino)ethoxy]phenyl]benzoyl as the hydrochloride salt was also effective in the treatment of PCP. Aerosol prophylaxis (two 60-minute treatments twice a week for 5 weeks) was highly effective. in preventing PCP in the infected immune suppressed rats. Aerosol therapy with 5, 10, 25, or 50 mg/ml of aerosolized solution reduced the number of cysts in the lungs by >99% when compared to controls. Similar results were obtained by i.p. dosage.

The following examples of compounds of the invention and the manner of their preparation further describe the present invention.

N-Acylation of Cyclohexpeptide Nuclei

The preparation of the derivatives of the A30912A nucleus was accomplished by the following general procedure, with Table 27 listing these derivatives.

The A30912A nucleus and the 2,4,5-trichlorophenol ester are dissolved in dimethylformamide (25-50 ml) and stirred for 17-65 hours at room temperature. The solvent is removed *in vacuo* and the residue is slurried in ether and collected by filtration. The solid product is washed with methylene chloride and then dissolved in either methanol or acetonitrile/water (1:1 v/v). This solution is injected on a Waters 600E semi-preparative chromatography system using a Rainin Dynamax-60A C₁₈ reverse-phase column. The column is eluted beginning with 20-40% aqueous acetonitrile and 0.5% monobasic ammonium phosphate (w/v) (monitored by UV at 230 nm and at a flow rate of 20 ml/min) until the unreacted A30912A nucleus is eluted and then deleting the buffer and eluting the product peak in aqueous acetonitrile. The fraction containing the product is evaporated *in vacuo* or lyophilized to provide the pure compound. The product may be analyzed by the same HPLC instrument using a Waters C₁₈ Micro Bondapak column and eluting with 40% aqueous acetonitrile containing 0.5% monobasic ammonium phosphate (w/v) at a 2 ml/min flow rate and monitoring the UV at 230 nm. The products may also be analyzed by fast atom bombardment mass spectrometry (FABMS). (In the compounds used, R'=R"=CH₃, R=OH, R^y=OH, R₁=H, R₇=OH, and R₂ is as defined).

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5	H ₃ C(CH ₂₎₅ N	CH ₂ C	H ₃ C(CH ₂) ₂	(CH ₂) ₂ 0-	CH ₃ (CH ₂) ₅ O	(H ₃ CCH ₂) ₂ CHCH ₂ O-	(CH ₃) ₃ C(CH ₂) ₂ O	СH ₂ 0⟨-	CH3(CH2),O	H ₃ C(CH ₂) ₄ O(CH ₂) ₂ O-	(H₃C) ₂ CH(CH ₂) ₂ O	Сн₂о	H ₃ C(CH ₂) ₃ O	
	Yook Y	N 20 0 1	N > 0			HCH ₂ O				(CH ₂) ₂ O	H ₂) ₂ 0			R ₂
15 20		14		•	ا چ	3 0 . 5	50	5	2	6	5	ن م	(A	Este Reactant
25	1000	1490	683	629	596	596	596	594	289	634	579	576	561	(mg) }
30	1.2	2.0	1.0	1.0	1.0	1.0	1.0	1.0	0.5	1.0	1.0	1.0		A30912A Pro Nucleus (g)
35	194 11	116 11	384 11	180 11	301 1	359 1	270	295	8	359 1	355	294 1	235	Product (mg) F
40	1190*+ 2	1195** 2	1147**	1104**	1100+ 10	1100* 9	1100* 8	1098+	1083*	1130+	1086*	1062*	1072*	FABMS HPLO
45	2.41	2.06	1.92		10.24	9.13	8.15	6.44	6.08	5.79	5.75	4.46		HPLC

TABLE 27

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10 15	H3C(CH2)50	H3C(CH2)30		$H_3C(CH_2)_7 \longrightarrow \begin{pmatrix} & & & \\ & & & \end{pmatrix} \stackrel{\Omega}{\longrightarrow} $		$H_3C(CH_2)_5 = \left\langle \begin{array}{c} \\ \\ \end{array} \right\rangle$	н ₃ с(сн ₂₎₃ ()-сн ₂ о ()-	H ₃ C(CH ₂) ₉ O(CH ₂) ₂ O	H ₃ C(CH ₂),O(CH ₂) ₂ O	H ₃ C(CH ₂) ₆ O(CH ₂) ₂ O	$H_3C(CH_2)_3$		OcH2 NOOF JE		OCH2N YOUNG	R ₂	
20	616	291	501	546	514	511	579	313	593	287	571	596	750	810	734	Ester Reactant (mg)	TABL
25	1.0	0.5	1.0	1.0	1.0	1.0	1.0	0.5	1.0	0.5	1.0	1.0	1.0	1.0	0.9	A30912A Nucleus (g)	TABLE 27 continued
30	341	98	218	285	287	322	293	. 104	307	110	295	190	126	230	303	Product (mg) FABMS	ъ.
35	1116*	1088*	1002**	1060+	1034*	1032+	1086*	1124+	1096*	1082*	1058++	1078**	1201**	1187**	1202*		
40	11.56	3.96	2.53	12.48	6.14	5.10	6.14	19.04	7.28	4.52	7.91	6.30	3.50	2.52	2.21	HPLC Retention (min)	
45																	

5	(m-1)+ Na +: ** (m+1); ***		1 ₃ C(CH ₂),	R2	
15 20	* m+[Na]+	566	534	Ester Reactant (mg)	TABLE
25		1.0	1.0	A30912A Nucleus (g)	E 27 continued
30		. 8	215	Product (mg) FABMS	<u> </u>
35		1054**	1050***		
40		3.89	7.59	HPLC Retention (min)	
45				e	

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Compounds such as those listed in Table 27 could be further modified at the phenolic hydroxy to provide R7 = -OPO₃HNa as shown in Table 28. The procedure is as follows:

The lipopeptide (1 equivalent) and tetrabenzylpyrophosphate (2 equivalents) were dissolved in dimethyl-

formamide which had been dri d ov r 13X molecular sieves. Lithium hydroxide monohydrate (5 equival nts) was added and the stirred solution was monitored by HPLC. After 0.5 hr and 1 hr more lithium hydroxid (5 quival nts) was added. Betw n 1 and 2 hrs. the reaction was quench d with glacial acetic acid, the solvent remov d under vacuum, and the residue purified ov reasemi-preparative C18 reverse-phase column using an aqueous acetonitrile eluent. The purified product was dissolved in (1/1) acetic acid/water with sodium acetate (1 equivalent) and 10% Pd/C catalyst. The solution was placed under an atmosphere of hydrogen gas and

stirred for 1 hr. After filtering to remove the catalyst, the solution was lyophilized to provide the pure final product. The purity was assessed by analytical HPLC and the product was analyzed by fast atom bombardment mass spectrometry (FABMS).

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		IABLE 28	82		
R ₂	Start. Mat. R ₇	Wt. (mg)	Prod.	Wt. (mg)	FABMS
(CH ₂) ₂ () 2	-0H	500	-OPO ₃ HNa	140	1184
H3C(CH2)30()	-011	300	-OPO3HNa	62	1228.4472*
* 11)+1					

Preparation f dideoxy cyclohexapeptide

The preparation of the dideoxy compounds may be accomplished by the following procedure with Table 29 listing derivatives.

To a suspension of a non-dideoxy cyclohexapeptide (formula (I) where R=OH and R_2 is hydrogen or acyl), in dichloromethane is added the reducing agent triethylsilane in dichloromethane. The solution is stirred and the volatile components are removed under reduced pressure and the residue triturated with diethyl ether. The compound is purified using HPLC, and the product lyophilized.

10 Example

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Dideoxycilofungin

To a suspension of cilofungin (10.00 g, 9.71 mmol) in dichloromethane (100 ml) was added a solution of triethylsilane (96 m1, 602 mmol) in dichloromethane (50 ml). Trifluoroacetic acid (46.4 ml, 602 mmol) was added as a solution in dichloromethane (50 ml) over 15 minutes. The solution was stirred at room temperature for two hours. The volatile reaction components were removed under reduced pressure and the residue triturated with diethyl ether. The compound was purified by reversed phase HPLC by means of a "Prep LC/System 500" unit (Waters Associates, Inc., Milford, Mass.) using a Prep Pak 500/C₁₈ Column (Waters Associates, Inc.) as the stationary phase. The column eluted with a gradient mobile phase using CH₃CN/H₂O (10:90 to 20:80 v/v) at 500 psi. The product containing fractions were pooled, evaporated under reduced pressure, and lyophilized from p-dioxane to yield dideoxycilofungin (6.66 g, 68.7%). FAB-MS: m/z calc. for C₄₉H₇₂N₇O₁₅, 998.5086; found, 998.512; UVλ(EtOH)nm(ε) 202.60(61012), 256.20(18569).

Table 29, indicates R_2 , the amount of the cyclic hexapeptide and reagents, and yield of dideoxy compounds prepared as described above. (R'=R"=R"=CH₃, R₁=H and R=R^Y=R₇=OH); T.E.S. = triethylsilane; TFA=trifluoroacetic acid; numbers are weights in grams).

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5 10	О−С6H13	io .HCI	Ö,C10H21	C ₁₂ H ₂₅	(C ₁₀ H ₂₀) -0	F ₂
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25 30	0.500	2.00	0.500	0.500	0.500	Tab Starting Material
35	3.50	9.49	2.63	2.47	0.256	Table 29
40	3.44	9.72	2.57	2.42	0.251	TEA
45	0.291	1.47	0.392	0.063	0.095	Yield

A compound of the formula

the preparation of which is discussed just prior to Table 27, can also be further modified at the phenolic hydroxy to provide R_7 =-OPO₃HNa, as indicated in the two paragraphs prior to Table 28. The compound produced is as follows:

The product was analyzed by FABMS (using Lit) to give a peak at 1226.4853 (calculated for C₅₈H₇₄N₇O₂₀PLi=1226.4886). Also, when analyzed by HPLC using a C18 reverse-phase column and eluting with 55% aqueous acetonitrile with 0.5% acetic acid at 2 ml/min and monitoring by UV at 280 nm, the compound had a ret ntion tim of 1.72 min.

Claims

1. A compound of the formula (1):

wherein

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R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" and R" are independently methyl or hydrogen;

R and RY are independently hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy; and

I) R₂ is a substituted benzoyl group represented by the formula

wherein

A) R₃ is a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m and n are integers of from 2 to 4, and p is 0 or 1; or

B) R₃ is an unsaturated hydrocarbon group represented by the formula

wherein Y is -C≡C- or -CH=CH-; or

C) R_3 is a group of the formula -O-(CH₂)_m-G, wherein m is as defined and G is C_7 - C_{10} bicycloalkyl

or C₇-C₁₄ tricycloalkyl; or

D) R₃ is quinolyl; or

II) R2 is an acyl group represented by the formula

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wh rein

Z is -O-, -C≡C-, -CH=CH-, -CH₂-CH₂-, -CH₂-, or a carbon to carbon bond;

A) R_4 is hydrogen, C_2 - C_{12} alkynyl, C_2 - C_{12} substituted alkynyl, C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_1 - C_{12} alkoxy, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, qui-

nolyl or phenyl; or

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B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkenyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, phenyl, substituted phenyl, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or

C) R4 is phenyl substituted with C1-C8 alkoxy substituted by fluoro, bromo, chloro or iodo; or

D) R_4 is C_1 - C_{12} alkoxy substituted with C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_2 - C_{12} alkynyl, amino, C_1 - C_4 alkylamino, di- $(C_1$ - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or

E) R₄ is C₁-C₁₂ alkoxy substituted with a group of the formula

O || -NHCR₈

wherein R₈ is C₁-C₆ alkoxy optionally substituted with phenyl; or

F) R4 is a group represented by the formula

wherein p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or piperazino, and R_5 is hydrogen, C_1 - C_{12} alkyl, C_3 - C_{12} cycloalkyl, benzyl or C_3 - C_{12} cycloalkylmethyl; or

G) R₄ is a group represented by the formula

wherein Y has the same meanings defined above; and

 R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} substituted alkyl; C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzothiazolyl, thienyl, phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(CH_2)p'-W- R_5 , or C_1 - C_6 alkoxy substituted by fluoro, bromo, iodo or chloro; or

R₆ is a phenyl substituted by a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined above; or

III) R₂ is a group having the formula

-CHNNNR*

wherein R^x is C₁-C₁₂ alkoxy or a polyoxa-alkyl group represented by the formula $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12})$ alkyl)

wherein m,n and p are as defined above; or

IV) R₂ is a group having the formula

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$$-C - (C_1 - C_{12} \text{ alkyl}) - O - C$$

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wherein R₉ is phenyl, C₁-C₁₂ alkyl, or C₁-C₁₂ alkoxy; or

V) R_2 is naphthoyl substituted with R_4 ; and the pharmaceutically acceptable non-toxic salts thereof; with the proviso that when

R' is methyl or NH₂C(O)CH₂-;

R" is methyl;

R" is methyl;

RY is hydroxy;

R is hydroxy; and

either a) or b):

- a) R₁ is hydroxysulfonyloxy and R₇ is hydroxy, hydroxysulfonyloxy or phosphonooxy;
- b) R_1 is hydrog $\,$ n or hydroxysulfonyloxy and R_7 is hydroxysulfonyloxy or phosphonooxy; R_2 is not

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i)

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wherein R₃ is

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein p=O; nor ii)

wherein Z is a carbon to carbon bond or -O- and R₄ is C₁-C₁₂ alkoxy; nor iii) naphthoyl substituted by R₄ wherein R₄ is hydrogen, phenyl, or C₁-C₁₂ alkoxy.

A compound of the formula (1): 2.

H 25 N- R2 Ry HN *30* R' -R" NH ОН OH (1) 0 35 Ry 40

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wherein

R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" and R" are independently methyl or hydrogen;

R and RY are independently hydroxy or hydrogen;

R₁ is hydroxy or hydrogen;

R₇ is hydroxy or hydrogen; and

I) R₂ is a substituted benzoyl group represented by the formula

wherein

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A) R₃ is a polyoxa-alkyl group repres nted by the formula

 $-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$

wherein m and n are integers of from 2 to 4, and p is 0 or 1; or

B) R₃ is an unsaturated hydrocarbon group represented by the formula

wherein Y is -C≡C- or -CH=CH-; or

C) R_3 is a group of the formula -O-(CH₂)_m-G, wherein m is as defined and G is C_7 - C_{10} bicycloalkyl or C_7 - C_{14} tricycloalkyl; or

D) R₃ is quinolyl; or

II) R2 is an acyl group represented by the formula

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wherein

Z is -O-, -C=C-, -CH=CH-, -CH₂-CH₂-, -CH₂-, or a carbon to carbon bond;

A) R_4 is hydrogen, C_2 - C_{12} alkynyl, C_2 - C_{12} substituted alkynyl, C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_1 - C_{12} alkoxy, C_3 - C_{12} cycloalkoxy, naphthyl, pyridyl, thienyl, benzothienyl, quinolyl or phenyl; or

B) R_4 is phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_2 - C_{12} alkenyl, C_2 - C_{12} alkynyl, C_1 - C_{12} substituted alkyl, C_2 - C_{12} substituted alkenyl, C_2 - C_{12} substituted alkynyl, C_1 - C_{12} alkonyl, trifluoromethyl, phenyl, substituted phenyl, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or

C) R₄ is phenyl substituted with C₁-C₆ alkoxy substituted by fluoro, bromo, chloro or iodo; or

D) R_4 is C_1 - C_{12} alkoxy substituted with C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, C_2 - C_{12} alkynyl, amino, C_1 - C_4 alkylamino, di- $(C_1$ - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino, phenyl substituted with a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m,n and p are as defined; or

E) R₄ is C₁-C₁₂ alkoxy substituted with a group of the formula



wherein R₈ is C₁-C₆ alkoxy optionally substituted with phenyl; or

F) R₄ is a group represented by the formula

wherein p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or piperazino, and R_5 is hydrogen, C_1 - C_{12} alkyl, C_3 - C_{12} cycloalkyl, benzyl or C_3 - C_{12} cycloalkylmethyl; or

G) R₄ is a group represented by the formula

$$-Y-R_{\theta}$$

wherein Y has the same meanings defined above; and

R₆ is C₁-C₁₂ alkyl, C₁-C₁₂ substituted alkyl; C₃-C₁₂ cycloalkyl, C₇-C₁₀ bicycloalkyl, C₇-C₁₄ tricycloalkyl, phenyl, C₃-C₁₂ cycloalkenyl, naphthyl, benzothiazolyl, thienyl, phenyl substituted by amino, C₁-C₁₂ alkylthio, halogen, C₁-C₁₂ alkyl, C₂-C₁₂ alkenyl, C₂-C₁₂ alkynyl, C₁-C₁₂ alkoxy, trifluoromethyl, -O-(CH₂)p'-W-R₅, or C₁-C₆ alkoxy substitut d by fluoro, bromo, iodo or chloro; or

R₈ is a ph nyl substitut d by a polyoxa-alkyl group repres nted by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wh rein m,n and p are as defined above; or

III) R₂ is a group having the formula

wherein R^x is C_1 - C_{12} alkoxy or a polyoxa-alkyl group represented by the formula $-O-(CH_2)_m$ - $[O-(CH_2)_n]_p$ - $O-(C_1-C_{12}$ alkyl)

wherein m,n and p are as defined above; or IV) R_2 is a group having the formula

$$-C - (C_1-C_{12} \text{ alkyl})-O$$

wherein R₉ is phenyl, C₁-C₁₂ alkyl, or C₁-C₁₂ alkoxy; or

V) R₂ is naphthoyl substituted with R₄; and the pharmaceutically acceptable non-toxic salts thereof.

- 3. A compound as recited in Claims 1 or 2 wherein R', R' and R'' are methyl, R₁ is hydrogen, and R₇ and R⁴ are OH.
- 4. A compound as recited in Claims 1 or 2 wherein R₂ is of the formula

wherein Z is a carbon to carbon bond; and

R₄ is C₁-C₁₂ alkoxy, C₃-C₇ cycloalkoxy, C₁-C₈ alkoxy substituted by C₃-C₇ cycloalkyl; or

R₄ is phenyl substituted by C₁-C₁₂ alkoxy or phenyl substituted with a polyoxa-alkyl group of the formula

 $-O-(CH_2)_m-[O-(CH_2)_n]P-O-(C_1-C_{12} \text{ alkyl}); \text{ or }$

 R_4 is a group of the formula -Y-R₆, wherein Y is an acetylenic bond and R₆ is C_1 -C₆ alkyl, phenyl, or phenyl substituted with a polyoxa-alkyl group of the formula

-O-(CH₂)_m-[O-(CH₂)n]P-O-(C₁-C₁₂ alkyl).

5. A compound as recited in claims 1 or 2 wherein R₂ is of the formula

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wherein Z is -C≡C-; and

R₄ is phenyl substituted by C₁-C₁₂ alkoxy or phenyl substituted with a polyoxa-alkyl group of the formula

 $-O-(CH_2)_m-[O-(CH_2)_n]P-O-(C_1-C_{12} \text{ alkyl})$

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6. A compound as recited in Claims 1 or 2 wherein R₂ is of the formula

$$-C(O)$$
 $-Z$ $-Z$

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wherein Z is a carbon to carbon bond and R₄ is a group of the formula -O-(CH₂)_p-W-R₅

wherein W is a piperidine group.

- 7. A compound as recited in Claims 1 or 2 wherein R is hydrogen.
- 8. A compound as recited in claims 1 or 2 wherein R₂ is 4-(4-n-hexyloxyphenyl)benzoyl,4-(4-n-heptyloxyphenyl)benzoyl, 4-(4-n-octyloxyphenyl)benzoyl, 4-[4-(3,3-dimethylbutoxy)phenyl]benzoyl, 4-[4-(2-cycloperyloxyethoxy)phenyl]benzoyl, 4-[4-(phenylethynyl)phenyl]benzoyl, 4-[4-(n-butylethynyl)phenyl]benzoyl, or 4-[4-[2-(4-cycloperyloxyethoxy)phenyl]benzoyl, 2-[4-(n-butylethynyl)phenyl]benzoyl, 3-[4-(n-butylethynyl)phenyl]benzoyl, 3-[4-(n-butylethynyl)p
 - 9. A compound of the formula (1):

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$$R''$$
 R''
 wherein

R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" and R" are independently methyl or hydrogen;

R and RY are independently hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy; and

I) R₂ is a group of the formula

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 H ₃ C(CH ₂) ₂ O
(H ₃ C) ₃ CO(CH ₂) ₂ O
H ₃ C(CH ₂) ₃ O(CH ₂) ₂ O
$H_3C(CH_2)_2O$
H ₃ C(CH ₂) ₃ O(CH ₂) ₂ O
(H ₃ C) ₃ CO(CH ₂) ₂ O
H ₃ C(CH ₂) ₃ O
H ₃ C(CH ₂) ₄ O
H ₃ C(CH ₂) ₅ O
H ₃ C(CH ₂) ₃ O(CH ₂) ₂ O
(H ₃ C) ₃ CO(CH ₂) ₂ O

R', R'' and R''' are methyl, R_1 is hydrogen and R_7 and R^Y are hydroxy and pharmaceutically acceptable salts thereof.

10. A compound of the formula (1):

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$$H_3C$$
 H_3C
 H_3C
 H_4
 H_5
 H_5
 H_7
 H

wherein R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" is methyl or hydrogen;

R is hydroxy or hydrogen;

R₁ is hydroxy, hydrogen, or hydroxysulfonyloxy;

R₇ is hydroxy, hydrogen, hydroxysulfonyloxy or phosphonooxy;

R₂ is a substituted benzoyl group represented by the formula

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wherein R₃ is a polyoxa-alkyl group represented by the formula

$$-O-(CH_2)_m-[O-(CH_2)_n]_p-O-(C_1-C_{12} \text{ alkyl})$$

wherein m and n are integers of from 2 to 4, and p is 0 or 1; or R_3 is an unsaturated hydrocarbon group represented by the formula

wherein Y is -C≡C- or -CH=CH-;

or R_3 is a group of the formula -O-(CH₂)_m-G, wherein m is as defined and G is C_7 - C_{10} bicycloalkyl or C_{7} - C_{14} tricycloalkyl;

or R2 is an acyl group represented by the formula

wherein Z is -O-, -C≡C-, -CH=CH-, -CH₂-CH₂-, or a carbon to carbon bond;

 R_4 is hydrogen, C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, phenyl, phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_1 - C_{12} alkoxy, trifluoromethyl, ph. nyl, or C_1 - C_8 alkoxy substituted by fluoro, bromo, chioro or iodo;

or R_4 is C_1 - C_{12} alkoxy, C_3 - C_{12} cycloalkoxy, C_1 - C_{12} alkoxy substituted by C_3 - C_{12} cycloalkyl, C_7 - C_{10} bicycloalkyl, C_7 - C_{14} tricycloalkyl, amino, C_1 - C_4 alkylamino, di-(C_1 - C_4 alkyl)amino, C_1 - C_{12} alkanoylamino or a group of the formula

O || -NHCR₈

wherein R_8 is C_1 - C_6 alkoxy optionally substituted with phenyl; or R_4 is a group represented by the formula -O- $(CH_2)_{p'}$ -W- R_5

wherein p' is an integer of from 2 to 4; W is pyrrolidino, piperidino or piperazino, and R_5 is hydrogen, C_{12} alkyl, C_3 - C_{12} cycloalkyl, benzyl or C_3 - C_{12} cycloalkylmethyl;

or R_4 is a group represented by the formula -Y- R_6 wherein Y has the same meanings defined above and R_6 is C_1 - C_{12} alkyl, C_1 - C_{12} alkyl substituted by phenyl; C_3 - C_{12} cycloalkyl, phenyl, C_3 - C_{12} cycloalkenyl, naphthyl, benzthiazol-2-yl, or phenyl substituted by amino, C_1 - C_{12} alkylthio, halogen, C_1 - C_{12} alkyl, C_1 - C_{12} alkenyl, C_1 - C_{12} alkoxy, trifluoromethyl, -O-(CH₂)p'-W- R_6 , or C_1 - C_6 alkoxy substituted by fluoro, bromo, iodo or chloro; or

R₂ is a group selected from

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$$C = C - (C_1 - C_{12} \text{ alkyl}) - C - C_{12}$$

wherein R₉ is phenyl, C₁-C₁₂ alkyl, or C₁-C₁₂ alkoxy; or

 R_2 is naphthoyl substituted with R_4 ; and the pharmaceutically acceptable non-toxic salts thereof; with the proviso that when

R' is methyl or NH₂C(O)CH₂-;

R" is methyl;

R is hydroxy; and

either

a) R₁ is hydroxysulfonyloxy and R₇ is hydroxy, hydroxysulfonyloxy or phosphonooxy; or

b) R_1 is hydrogen or hydroxysulfonyloxy and R_7 is hydroxysulfonyloxy or phosphonooxy; R_2 is not

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wherein R_3 is -O-(CH₂)_m-[O-(CH₂)_n]_p-O-(C₁-C₁₂ alkyl) wherein p=O; nor ii)

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wherein Z is a carbon to carbon bond or -O- and R_4 is C_1 - C_{12} alkoxy; nor iii) naphthoyl substituted by R_4 wherein R_4 is hydrogen, phenyl, or C_1 - C_{12} alkoxy.

- 11. A compound as recited in claim 10 wherein R₁ is not hydroxysulfonyloxy and R₇ is not hydroxysulfonyloxy or phosphonoxy.
- 12. A compound of any of claims 1-11 for use in inhibiting parasitic activity.
 - 13. A compound of claims 1-11 for use in inhibiting fungal activity.
- 14. A compound of any of claims 1-11 for use in inhibiting the growth of organisms responsible for opportunistic infections in immunosuppressed individuals.
 - 15. A compound of claims 1-11 for use in inhibiting the growth of Pneumocystis carinii.
- 16. A pharmaceutical formulation comprising a compound of any of Claims 1-11 and a suitable pharmaceutical carrier.
 - 17. A process for the preparation of a compound of the formula (1):

wherein R' is hydrogen, methyl or NH₂C(O)CH₂-;

R" and R" is methyl or hydrogen;

R is hydrogen;

RY is hydroxy or hydrogen,

R₁ is hydroxy, or hydrogen;

R₇ is hydroxy, or hydrogen; and

R₂ is hydrogen or acyl;

comprising the step of subjecting a compound of formula (I) wherein R=OH, to a strong acid in the presence of a reducing agent, in a suitable solvent.

18. A compound of the formula

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EUROPEAN SEARCH REPORT

Application Number

	OCCIVIENTS CONSIDE	RED TO BE RELEVA	1 1 1	EP 93302064.6
Category	Citation of document with indicate of relevant passage		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.5)
A	EP - A - 0 448 34 (MERCK & CO. INC * Claims 1-7	•)	1-18	C 07 K 7/56 A 61 K 37/02
A	EP - A - 0 448 39 (MERCK & CO. INC * Claims 1-7	.)	1-18	
A	EP - A - 0 447 18 (MERCK & CO. INC. * Claims 1-7		1-18	
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				TECHNICAL FIELDS SEARCHED (Int. CL5)
				C 07 K 7/00 A 61 K 37/00
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	The present search report has been d	rawn up for all claims		
	Place of search VIENNA	Date of completion of the search 28-05-1993	S	Examiner CHARF
X : partic Y : partic docum	ATEGORY OF CITED DOCUMENTS cularly relevant if taken alone cularly relevant if combined with another ment of the same category cological background	E : earlier patent after the filin D : document cit	ciple underlying the document, but public date application of for other reasons	ished on, or